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(54) Title: PARATHYROID HORMONE RECEPTOR AND DNA ENCODING SAME

(57) Abstract

DNA encoding a parathyroid hormone receptor; production and isolation of recombinant and synthetic parathyroid hormone receptor polypeptides and fragments; antibodies to parathyroid hormone receptors and receptor fragments; methods for screening candidate compounds for antagonistic or agonistic effects on parathyroid hormone receptor action; and diagnostic and therapeutic methods of these compounds are disclosed.

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PARATHYROID HORMONE RECEPTOR AND DNA ENCODING SAME Background of the Invention

Partial funding of the work described herein was 5 provided by the U.S. Government, which has certain rights to the invention.

The invention relates to endocrine receptors.

A crucial step in the expression of hormonal action is the interaction of hormones with receptors on the plasma membrane surface of target cells. The formation of hormone-receptor complexes allows the transduction of extracellular signals into the cell to elicit a variety of biological responses. For example, binding of a hormone such as follicle stimulating hormone

- 15 (FSH), luteinizing hormone (LH), thyroid stimulating hormone (TSH), and chorionic gonadotropin (CG), to its cell surface receptor induces a conformational change in the receptor, resulting in the association of the receptor with a transductor molecule, the stimulatory
- guanine nucleotide (GTP) binding protein, a component of which is (G_s) . This association stimulates adenylate cyclase activity which in turn triggers other cellular processes such as protein phosphorylation, steroid synthesis and secretion, and the modulation of ion flux.
- Binding of other hormones, including arginine vasopressin (VP), angiotensin II, and norepinephrine, to their cell surface receptors results in the activation of other types of GTP binding proteins components such as (G_p) , which in turn stimulates the activity of the enzyme
- 30 phospholipase C. The products of phospholipase C hydrolysis initiate a complex cascade of cellular events, including the mobilization of intracellular calcium and protein phosphorylation.

Parathyroid hormone (PTH) is a major regulator of 35 calcium homeostasis whose principal target cells occur in

bone and kidney. Regulation of calcium concentration is necessary for the normal function of the gastrointestinal, skeletal, neurologic, neuromuscular, and cardiovascular systems. PTH synthesis and release 5 are controlled principally by the serum calcium level: a low level stimulates and a high level suppresses both the hormone synthesis and release. PTH, in turn, maintains the serum calcium level by directly or indirectly promoting calcium entry into the blood at three sites of 10 calcium exchange: qut, bone and kidney. PTH contributes to net gastrointestinal absorption of calcium by favoring the renal synthesis of the active form of vitamin D. promotes calcium resorption from bone by inhibiting osteoblasts and, indirectly, by stimulating 15 differentiation of the bone-resorbing cells, osteoclasts. It also mediates at least three main effects on the kidney: stimulation of tubular calcium reabsorption, enhancement of phosphate clearance, and promotion of an increase in the enzyme that completes synthesis of the 20 active form of vitamin D. PTH exerts these effects primarily through receptor-mediated activation of adenylate cyclase, although receptor-mediated activation of phospholipase C by PTH has also been reported (Hruska

Disruption of calcium homeostasis may produce many clinical disorders (e.g., severe bone disease, anemia, renal impairment, ulcers, myopathy, and neuropathy) and usually results from conditions which produce an alteration in the level of parathyroid hormone.

et al., J. Clin. Invest. 79:230, 1987).

30 Hypercalcemia is a condition which is characterized by an elevation in the serum calcium level. It is often associated with primary hyperparathyroidism in which an excess of PTH production occurs as a result of a lesion (e.g., adenoma, hyperplasia or carcinoma) of the

35 parathyroid glands. Another type of hypercalcemia,

humoral hypercalcemia of malignancy (HHM), is the most common paraneoplastic syndrome. It appears to result in most instances from the production by tumors (e.g., squamous, renal, ovarian or bladder carcinomas) of a 5 novel class of protein hormone which shares amino acid homology with PTH. These PTH-related proteins (PTHrP) appear to mimic certain of the renal and skeletal actions of PTH and are believed to interact with the PTH receptor in these tissues. PTHrP is normally found at low levels 10 in many tissues, including keratinocytes, brain, pituitary, parathyroid, adrenal cortex, medulla, fetal liver, osteoblast-like cells and lactating mammary tissues. In many HHM malignancies, PTHrP is found in the circulatory system at high levels, thereby producing the 15 elevated calcium levels associated with HHM.

Summary of the Invention

The invention features isolated DNA comprising a DNA sequence encoding a cell receptor, preferably a parathyroid hormone receptor, of a vertebrate animal, 20 which receptor has an amino acid sequence with at least 30% (preferably at least 50%, even more preferably at least 60%, and most preferably at least 75%) identity to the amino acid sequence shown in FIG. 3 (SEQ ID NO.: 3): i.e., when the closest match is made between the two 25 amino acid sequences (using standard methods), at least 30% of the amino acid residues of the former sequence are identical to the amino acid residues of the latter sequence. By "isolated" is meant that the DNA is free of the coding sequences of those genes that, in the 30 naturally-occurring genome of the organism (if any) from which the DNA of the invention is derived, immediately flank the gene encoding the DNA of the invention. isolated DNA may be single-stranded or double-stranded, and may be genomic DNA, cDNA, recombinant hybrid DNA, or

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synthetic DNA. It may be identical to a naturallyoccurring, cell receptor- (e.g. PTH receptor) encoding DNA sequence, or may differ from such sequence by the deletion, addition, or substitution of one or more 5 nucleotides. Single-stranded DNAs of the invention are generally at least 8 nucleotides long, (preferably at least 18 nucleotides long, and more preferably at least 30 nucleotides long) ranging up to full length of the gene or cDNA; they preferably are detectably labelled for 10 use as hybridization probes, and may be antisense. Preferably, the isolated DNA hybridizes under conditions of high stringency to all or part of the DNA sequence show in FIG. 1 (SEQ ID NO.:1), FIG. 2 (SEQ ID NO.:2), FIG. 3 (SEQ ID NO.:3), or FIG. 6 (SEQ ID NO.:4). By 15 "high stringency" is meant, for example, conditions such as those described herein below for the isolation of human kidney PTH receptor cDNA (also see Current Protocols in Molecular Biology, John Wiley & Sons, New York, 1989, hereby incorporated by reference). Most 20 preferably, the animal is a mammal (such as an opossum, a rat, or a human), and the DNA sequence encodes substantially all of the amino acid sequence shown in FIG. 1 (SEQ ID NO.:1), FIG. 2 (SEQ ID NO.:2), FIG. 3 (SEQ ID NO.:3) or FIG. 6 (SEQ ID NO.:4); or is encoded by the 25 coding sequence of one of the plasmids deposited with the American Type Culture Collection (ATCC) and designated ATCC Accession No. 68570 or 68571. The DNA of the invention may be incorporated into a vector [which may be provided as a purified preparation (e.g., a vector 30 separated from the mixture of vectors which make up a library)] containing a DNA sequence encoding a cell receptor of the invention (e.g. parathyroid hormone receptor) or fragment of the receptor, and a cell or essentially homogenous population of cells (e.g.,

35 prokaryotic cells, or eukaryotic cells such as mammalian

cells) which contain the vector (or the isolated DNA described above). By "essentially homogenous" is meant that at least 99% of the cells contain the vector of the invention (or the isolated DNA, as the case may be).

Preferably, this vector (e.g., R15B) is capable of directing expression of a parathyroid hormone receptor (for example, in a cell transfected or transformed with the vector).

In another aspect, the invention features a cell receptor, preferably parathyroid hormone receptor, (or an essentially purified preparation thereof) produced by expression of a recombinant DNA molecule encoding the cell receptor. An "essentially purified preparation" is one which is substantially free of the proteins and lipids with which it is naturally associated.

In a related aspect, the invention features a polypeptide which includes a fragment of a naturally-occurring cell receptor of the invention. Preferably, the polypeptide includes a fragment of a naturally-

- occurring parathyroid hormone receptor which is capable of binding parathyroid hormone or parathyroid hormone-related protein. In preferred embodiments, this fragment is at least six amino acids long, and has a sequence selected from the group including:
- 25 (a) TNETREREVFDRLGMIYTVG; (SEQ ID NO.: 5)
 - (b) YLYSGFTLDEAERLTEEEL; (SEQ ID NO.: 6)
 - (c) VTFFLYFLATNYYWILVEG; (SEQ ID NO.: 7)
 - (d) Y-RATLANTGCWDLSSGHKKWIIQVP; (SEQ. ID NO.: 8)
 - (e) PYTEYSGTLWQIQMHYEM; (SEQ ID NO.: 9)
 - (f) DDVFTKEEQIFLLHRAQA; (SEQ ID NO.: 10)
 - (g) FFRLHCTRNY; (SEQ ID NO.: 11)
 - (h) EKKYLWGFTL; (SEQ ID NO.: 12)
 - (i) VLATKLRETNAGRCDTRQQYRKLLK; or (SEQ ID NO. 13)
- (j) a fragment (i.e., a portion at least six 35 residues long, but less than all) or analog of (a) (i)

30

which is capable of binding parathyroid hormone or parathyroid hormone-related protein [wherein "analog" denotes a peptide having a sequence at least 50% (and preferably at least 70%) identical to the peptide of 5 which it is an analog]. Preferably, the polypeptide of the invention is produced by expression of a recombinant DNA molecule or is synthetic (i.e., assembled by chemical rather than biological means). The invention provides a method for producing such a polypeptide, which method includes providing a cell containing isolated DNA encoding a cell receptor of the invention or receptor fragment and culturing this cell under conditions which permit expression of a polypeptide from the isolated DNA.

The invention also features an antibody 15 (monoclonal or poylclonal), and a purified preparation of an antibody, which is capable of forming an immune complex with a cell receptor of the invention (preferably a parathyroid hormone receptor such as a human PTH receptor) such antibody being generated by using as 20 antigen either (1) a polypeptide that includes a fragment of the cell receptor of the invention, or (2) a cell receptor of the invention which is on the surface of a cell. This antibody is preferably capable of neutralizing (i.e., partially or completely inhibiting) a 25 biological activity of the cell receptor of the invention (i.e., a component of one of the cascades naturally triggered by the receptor when its ligand binds to it). In preferred embodiments, the antibody of the invention is capable of forming an immune complex with parathyroid 30 hormone receptor and is capable of neutralizing a biological activity of the PTH receptor (i.e. adenylate cyclase activation or phospholipase C stimulation)

Also within the invention is a therapeutic composition including, in a pharmaceutically-acceptable carrier, (a) a cell receptor of the invention, (b) a

polypeptide containing a fragment of the cell receptor of the invention, or (c) an antibody to a cell receptor of the invention. These therapeutic compositions provide a means for treating various disorders characterized by overstimulation of the cell receptors of the invention by their ligand. In preferred embodiments, the polypeptides of the invention include the PTH receptor, fragments of the PTH receptor and antibodies which form immune complexes with the PTH receptor. These polypeptides and antibodies are useful as diagnostics, for distinguishing those cases of hypercalcemia related to PTH or PTHrP from those which are not.

The nucleic acid probes of the invention enable one of ordinary skill in the art of genetic engineering to identify and clone cell receptor homologs or cell receptors from any species which are related to the cell receptors of the invention, expanding the usefulness of the sequences of the invention.

Other features and advantages of the invention 20 will be apparent from the following description of the preferred embodiments and from the claims.

<u>Detailed Description</u>

The drawings will first be briefly described. DRAWINGS

FIG. 1 is a representation of the nucleic acid and amino acid sequence encoding the opossum kidney PTH/PTHrP receptor clone, OK-H. (SEQ ID NO.: 1)

FIG. 2 is a representation of the nucleic acid and amino acid sequence encoding the opossum kidney PTH/PTHrP receptor clone, OK-O. (SEQ ID NO.: 2)

FIG. 3 is a representation of the nucleic acid and amino acid sequence encoding the rat bone PTH/PTHrP receptor clone, R15B. (SEQ ID NO.: 3)

- FIG. 4 is a comparison of the deduced amino acid sequences encoded by cDNAs from clones OK-O and R15B.
- FIG. 5 is a comparison of the deduced amino acid sequences of OK-O, OK-H and R15B, lined up according to sequence homology.
 - FIG. 6 is a representation of the nucleic acid and amino acid sequence encoding the human PTH/PTHrP receptor. (SEQ ID NO.: 4)
- FIG. 7 is a schematic representation of the rat bone PTH/PTHrP receptor cDNA, the human genomic DNA clone HPG1 and two cDNA clones encoding the human PTH/PTHrP receptor.
- FIG. 8 is a hydrophobicity plot of the deduced amino acid sequence of the human kidney PTH/PTHrP

 5 receptor. Predicted membrane-spanning domains I through
- 15 receptor. Predicted membrane-spanning domains I through VII are indicated; A, B and C indicate additional hydrophobic regions.
 - FIG. 9 is a graph illustrating binding of PTHrF to COS cells transfected with OK-H.
- FIG. 10 is a graph illustrating stimulation of intracellular free calcium by NlePTH in COS cells transfected with OK-H.
 - FIG. 11 is a graph illustrating binding of PTHrP to COS cells transfected with OK-O.
- FIG. 12 is a graph illustrating stimulation of intracellular free calcium by NlePTH in COS cells transfected with OK-O.
 - FIG. 13 is a graph illustrating binding of PTHrP to COS cells transfected with R15B.
- FIG. 14 is a graph illustrating stimulation of intracellular free calcium by NlePTH in COS cells transfected with R15B.
- FIG. 15 is a graph illustrating stimulation of inositol phosphate metabolism by NlePTH in COS cells transfected with OK-H, OK-O, or R15B.

FIG. 16 is a graph illustrating cyclic AMP accumulation in COS cells transfected with CDM-8, OK-H, R15B by NlePTH.

FIG. 17 are graphs illustrating binding of \$^{125}I^{-}\$

5 labelled PTH(1-34) (A and B) and \$^{125}I^{-}\$

36) (C and D) to COS-7 cells transiently expressing the human kidney (A and C) and the rat bone (B and D)

PTH/PTHrP receptor; competing ligands included PTH(1-34) (D), PTHrP(1-36) (*), PTH(3-34) (E), PTH(7-34) (+).

10 Data are given as % specific binding and represent the

mean±SD of at least three independent experiments.

FIG. 18 is a bar graph illustrating stimulated accumulation of intracellular cAMP in COS-7 cells transiently expressing the human kidney receptor. Data show the mean±SD, and are representative of at least three independent experiments.

FIG. 19 represents a Northern blot analysis of total RNA (~ 10 μ g/lane) prepared from human kidney (A) and SaOS-2 cells (B). The blot was hybridized with the full length cDNA encoding the human kidney PTH/PTHrP receptor; positions of 28S and 18S ribosomal RNA bands are indicated.

FIG. 20 represents a Southern blot analysis of human genomic DNA digested with SstI, HindIII, and XhoI
 25 (~10μg/lane. The blot was hybridized with the full length cDNA encoding the human kidney PTH/PTHrP receptor.

FIG. 21 is a schematic diagram of the proposed arrangement, in a cellular membrane, of PTH/PTHrP rat bone receptor encoded by R15B.

MATERIALS AND METHODS

GENERAL: [Nle^{8,18}, Tyr³⁴]bPTH(1-34)amide (PTH(1-34)),
[Nle^{8,18}, Tyr³⁴]bPTH(3-34)amide (PTH(3-34)), and [Nle^{8,18},
Tyr³⁴]bPTH(7-34)amide (PTH(7-34)) were obtained from
Bachem Fine Chemicals, Torrance, CA; [Tyr³⁶]PTHrP(1-

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- 36) amide (PTHrP(1-36)) was synthesized as described (Keutman et al., Endocrinology 117:1230, 1985) using an Applied Biosystems Synthesizer 420A. Dulbecco's modified Eagles medium (DMEM), EDTA/trypsin, and gentamycin were from GIBCO (Grand Island, NY); fetal bovine serum (FBS) was from Hiclone Laboratory, Logan, UT. Total RNA from human kidney was provided by Per Hellman, University Hospital, Uppsala, Sweden. Oligonucleotide primers were
- 10 Synthesizer. Restriction enzymes, Klenow enzyme, T4 polynucleotide Kinase and T4 DNA ligase were from New England Biolabs, Beverly, MA. Calf alkaline phosphatase was from Boehringer Mannheim, Germany. All other reagents were of highest purity available.

synthesized using an Applied Biosystems 380B DNA

15 CELLS

Cell lines used include COS cells, OK cells, SaOS-2 cells, CHO cells, AtT20 cells, LLC-PK1 cells, and UMR-106 cells, which are available from a variety of sources including the American Type Culture Collection (Rockland,

- 20 Maryland), Accession Nos. CRL1650, CRL6551, HTB85, CCL61, CCL89, CL101, and CRL1161, respectively. ROS 17/2 and ROS 17/2.8 are available from a number of sources including Dr. Gideon Rodan (Merck Laboratories, West Point, PA). MC-3T3 cells are derived from mouse bone
- 25 cells and are also available from a number of sources including Dr. Chohei Shigeno (Dept. of Biochem. Medicine, Hyoto Univ., Kyoto, Japan).

All cells were grown in a humidified 95% air, 5% CO₂ atmosphere and maintained in monolayer culture with

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F-12 or DMEM medium (Grand Island Biological Co.), supplemented with 5% or 10% fetal calf serum (M.A. Bioproducts, Walkersville, MD). The medium was changed every 3 or 4 days, and the cells were subcultured every 2 or 3 weeks by

trypsinization using standard methods. CLONING

Isolation of cDNA clones encoding the rat and opossum PTH/PTHrP receptors: Total RNA was initially isolated from rat osteosarcoma (ROS) cells (ROS 17/2.8) and opossum kidney (OK) cells, by standard methods using guanidium isothiocyanate (Ullrich et al., Science 196: 1313, 1977; Chirgwin et al. Biochemistry 24: 5294, 1979), and centrifugation through cesium chloride (Gilsen et

- al., Biochemistry 13: 2633, 1974). Poly A+ RNAs (mRNAs) were then recovered after passage of the total RNAs over oligo dT columns (Pharmacia, Piscataway, NJ) by the method of Aviv and Leder (Proc. Natl. Acad Sci. USA 69: 14087, 1972). The cDNA library from the ROS 17/2.8 mRNA
- was prepared from poly A+ RNA using the method of Gubler and Hoffman (Gene (Amst.) 25: 263, 1983). Oligo dT-primed and random-primed cDNAs were synthesized from poly A+ ROS 17/2.8 and OK cell mRNA, respectively (Aviv and Leder, supra). The cDNAs were ligated to BstX1 linkers
- 20 (Invitrogen, San Diego, CA) and size-selected by centrifugation (3 h, 55,000 xg) in a 5-20% potassium acetate gradient. The size-selected cDNA was then inserted into the plasmid vector, pcDNA I (Invitrogen), using the non-self annealing BstX1 restriction sites.
- 25 The resultant plasmid libraries were then used to transform

E. coli (MC1061/P3, Invitrogen) containing a larger helper plasmid, p3. The p3 plasmid possesses amber mutations in two genes which code for ampicillin and

- tetracycline resistance. Using ampicillin and tetracycline selection, only those cells containing both the p3 and a tRNA suppressor gene, which is contained within pcDNA I, were capable of growth. The transformed bacteria were then grown to confluence, and the plasmid
- 35 DNAs isolated using standard techniques (e.g., see

Ausebel et al., Current Protocols in Molecular Biology, John Wiley Sons, New York, 1989). These DNAs were then taken up in a DEAE-dextran solution, and used to transfect African Green Monkey kidney (COS) cells, which 5 had been grown to 75% confluence in "sideflasks" (Nunc, Denmark).

Screening for COS cells containing plasmids capable of expressing functionally-intact ROS or OK cell parathyroid hormone/parathyroid hormone related-protein 10 (PTH/PTHrP) receptor proteins was performed according to Gearing et al. (EMBO J. 8: 3676, 1989), with some minor modifications including DEAE-Dextran transfection in sideflasks. Forty-eight hours after transfection, the cells were tested for binding of 125 I-labeled [Tyr36]PTHrp 15 (1-36) amide, using methods previously described (Yamamoto et al., Endocrinology 122: 1208, 1988), with the following exceptions: the time and temperature of the incubation were 2h and room temperature, respectively. After rinsing, the cells were fixed with 1.25% 20 glutaraldehyde, and rinsed with 1% gelatin. snapping off the top of the sideflask, the remaining microscope slide was dipped into NTB-2 photographic emulsion (Eastman Kodak, Rochester, NY). After 3-4 days of exposure at 4°C, the slides were developed, fixed, and 25 stained with 0.03% toluene blue. Screening of each slide was performed under a light microscope (Olympus). pool of plasmid-DNA from ROS cells, and two pools of plasmid-DNA from OK cells, (10,000 independent clones), each gave rise to 3-4 transfected COS cells expressing 30 the PTH/PTHrP receptor. These pools were subsequently subdivided. The subpools were used to transfect COS cells, and single clones were identified that expressed receptor protein capable of binding the radioligand.

Isolation of cDNA and genomic DNA clones encoding
35 the human PTH/PTHrP receptor: A human kidney oligo dT-

primed cDNA library (1.7x106 independent clones) in lambda GT10 and a genomic library of human placental DNA (2.5x10⁶ independent clones) in EMBL3 (Sp6/T7) (Clontech, Palo Alto, CA) were screened by the plaque hybridization 5 technique (Sambrook et al., Molecular Cloning: A Laboratory Manual, 2nd Ed. pp. 108-113, Cold Spring Harbor Laboratory, Cold Spring Harbor, NY, 1989) with the 32P-labelled (random primed labelling kit Boehringer Mannheim, Germany) BamHI/NotI 1.8kb restriction enzyme 10 fragment encoding most of the coding sequence of the rat bone PTH/PTHrp receptor (Fig. 3). The nitrocellulose filters were incubated at 42°C for 4 hrs in a prehybridization solution containing 50% formamide, 4x saline sodium citrate (SSC; 1x SSC: 300 mM NaCl, 30 mM 15 NaCitrate, pH 7.0), 2x Denhardt's solution. 10% Dextran sulphate, 100 μ g/ml salmon sperm DNA (final

NaCitrate, pH 7.0), 2x Denhardt's solution, 10% Dextran sulphate, 100 μg/ml salmon sperm DNA (final concentration). The hybridizations were carried out in the same solution at 42°C for 18-24h. Filters were washed with 2x SSC/0.1% SDS for 30 minutes at room

20 temperature and then with ¹x SSC/0.1% SDS for 30 minutes at 45°C. The films were exposed at -80°C for 18-24h using intensifying screens.

About 1,000,000 clones were screened from each library. Positive clones were plaque-purified and lambda phage DNA was isolated (Sambrook et al., supra). Cloned inserts were removed from phage DNA by digestion with restriction endonucleases HindIII and EcoRI (lambda GT10 library), or with XhoI and SstI (EMBL3 library), and were then subcloned into pcDNAI (Invitrogen, San Diego, CA) using the appropriate, dephosphorylated restriction sites. Sequencing of the CsCl2-purified subclones was performed according to Sanger et al. (Biochem 74:5463, 1977) by the dideoxy termination method (Sequenase version 2 sequencing kit, United States Biochemical

35 Corporation, Cleveland, OH).

Reverse transcription and polymerase chain reaction (PCR): 3 μ g of poly (A)+ RNA from human kidney (Clontech, Palo Alto, CA) in 73.5 μ l of H₂O was incubated at 100°C for 30 seconds, quenched on ice, and then added to 20 μ l of 5x RT buffer (1x RT buffer: 40 mM Tris-HCl, pH 8.2, 40 mM KCl, 6.6 mM MgCl₂, 10 mM dithiothreitol, and dNTPs at 0.5 mM each), 2 μ l (4 units) RNasin (Promega Biotec, Madison, WI), 1 μ l (80 pmo/ μ l) of the human cDNA primer H12

- 10 (5'-AGATGAGGCTGTGCAGGT-3'; SEQ ID NO.: 14) and 80 units of avian myeloblastosis virus reverse transcriptase (Life Sciences, St. Petersburg, FL). The reaction mixture was incubated for 40 minutes at 42°C. One-tenth of the first strand synthesis reaction mixture was then amplified by
- 15 PCR in a final volume of 100 μ l containing 3 mM MgSO₄, 200 μ M dNTPs, 2 units of Vent polymerase (New England Biolab, Beverly, MA), and 2 μ M each of the forward and the reverse primers (PCR conditions: denaturing for 1 min at 94°C, annealing for 1 min at 50°C, and extension at 72°C for 3 minutes; 40 cycles).

Two independent PCRs were performed using two different forward primers: i) degenerate primer RK-1 (5'-GGAATTCCATGGGAGCGGCCCGGAT-3'; SEQ ID NO.: 15) based on

- the 5' coding end of the two previously cloned PTH/PTHrP receptors (described above), and ii) primer RK-2 (5'-CGGGATCCCGCGGCCCTAGGCGGT-3'; SEQ ID NO.: 16) based on the 5' untranslated region of the human genomic clone
- 30 HPG1. Both PCR reactions used the reverse primer H26 (5'AGTATAGCGTCCTTGACGA-3'; SEQ ID NO.: 17) representing nucleotides 713 to 731 of the coding region of the human PTH/PTHrP receptor (Fig. 4). PCR products were bluntended using Klenow enzyme and cloned into
- 35 dephosphorylated pcDNAI cut with EcoRV.

Northern blot analysis: Total RNA was extracted from SaOS-2 cells and from human kidney by the guanidine thiocyanate method (Chirgwin et al., Biochem. 18:5294, 1979). For Northern blot analysis, ~10 µg of total RNA was subjected to electrophoresis on a 1.5%/37% formaldehyde gel and blotted onto nitrocellulose filters (Schleicher and Schuell, Keene, NH). The hybridization conditions were the same as those for screening the phage libraries (see above). The filters were washed at a final stringency of 0.5x SSC/0.1% SDS for 30 min at 60°C and exposed for autoradiography.

Southern blot analysis: Human genomic DNA was prepared using the SDS/proteinase K method (Gross-Bellard et al., Eur. J. Biochem. 36:32, 1973). For Southern

15 analysis, ~10 μg of DNA was digested with SstI, PvuII and XhoI; subjected to electrophoresis on a 0.8% agarose gel; and blotted onto nitrocellulose membranes (Schleicher and Schuell, Keene, NH). The hybridization conditions were the same as those for screening the phage libraries (see above). The filters were washed at a final stringency of 0.5x SSC/0.1% SDS for 30 min at 55°C and exposed for autoradiography.

FUNCTIONAL ASSAYS

Tests to characterize the functional properties of the cloned receptors expressed on COS cells included:

- I) binding of PTH and PTHrP fragments and analogues, II) stimulation of cyclic AMP accumulation by PTH and PTHrP fragments and analogues,
- III) increase of intracellular free calcium by PTH 30 and PTHrP fragments and analogues, and
 - IV) activation of inositol phosphate metabolism by PTH and PTHrP fragments and analogues. The methodologies are as follows:

Radioreceptor Assay

[Nle⁸,Nle¹⁸,Tyr³⁴]bPTH-(1-34)amide (NlePTH), and [Tyr³⁶]PTHrP(1-36)amide(PTHrP) were iodinated with Na¹²⁵I (carrier free, New England Nuclear, Boston, MA) as 5 previously reported (Segre et al., J. Biol. Chem. 254: 6980, 1979), and purified by reverse-phase HPLC. brief, the labeled peptide was dissolved in 0.1% trifluoracetic acid (TFA), applied to a C18 Sep-pak cartridge (Waters Associates, Inc., Milford, MA) and 10 eluted with a solution of 60% acetonitrile in 0.1% TFA. After lyophilization, the radioligand then was applied to C_{18} - μ Bondapak column (3.9 mm x 30 cm. Waters Associates) and eluted over 30 min with a linear gradient of 30-50% acetonitrile-0.1% TFA at a flow rate of 2 ml/min. 15 radioligand eluted in two peaks; the first peak, which eluted at approximately 38% acetonitrile, was used in these studies because it gave higher total and specific .bindings. The specific activity was 500 ± 75 mCi/mg, which corresponds to an average iodine-peptide ratio of 20 1.

COS-7 cells were grown in 15 cm plates in DMEM, 10% heat-inactivated FBS, 10 mg/L gentamycin until 80-90% confluent. Twenty-four hours after transfection by the

- DEAE/Dextran method (Sambrook et al., supra), with 1-2 μ g of plasmid DNA, the cells were trypsinized and replated in multiwell plastic dishes (16 or 35 mm diameter, Costar, Cambridge, MA) at a cell concentration of 5 x 10⁴ cells/cm²). Cell number increased only slightly after
- 30 transfection. After continuing culture for another 48 h, radiorecepter assays were performed. The culture medium was replaced with buffer containing 50 mM Tris-HCL (pH 7.7),

100 mM NaCl, 2 mM CaCl₂, 5 mM KCL, 0.5% heat-inactivated 35 fetal bovine serum (GIBCO), and 5% heat-inactivated horse

serum (KC Biological Inc., Lenexa, KS) immediately before studies were initiated. Unless otherwise indicated, studies were conducted with cells incubated in this buffer at 15°C for 4 h with 4 x 10⁵ cpm/ml (9.6 x 10⁻¹¹ M) of ¹²⁵I-labeled NlePTH or PTHrP.

Incubations were terminated by aspirating the buffer, and repeatedly (x3) washing the culture dishes containing the adherent cells with chilled 0.9% NaCl solution, over a 15 sec period. Cell-bound radioactivity 10 was recovered by the sequential addition (x3) of 1 N NaOH (200 μl) to each well. After 30 min at room temperature, the NaOH was transferred to a glass tube. A second and third extraction with 1 N NaOH (200 μl) were combined with the first, and the total radioactivity was 15 counted in a γ-spectrometer (Packard Instruments, Downers Grove, IL). Tracer adherence to culture vessel without cells was negligible (<0.2% of total counts added), if vessels were preincubated with culture medium. Determinations of cAMP accumulation

Intracellular cAMP accumulation was measured as described previously (Abou-Samra et al., J. Biol. Chem. 262:1129, 1986). Cells in 24-well plates were rinsed with culture medium containing 0.1% BSA and 2mM IBMX. The cells were then incubated with PTH or PTHrP for 15 min. at 37° C. The supernatant was removed and the cells immediately frozen by placing the whole plate in dry ice powder. Intracellular cAMP was extracted by thawing the cells in 1ml of 50 mM HCl and analyzed by a specific radioimmunoassay using an anti-cAMP antibody (e.g.,

30 Sigma, St. Louis, MO). A cAMP analog (2'-O-monosuccinyl-adenosine 3':5'-cyclic monophosphate tyrosyl methyl ester, obtained from Sigma) which was used a tracer for cAMP was iodinated by the chloramine T method. Free iodine was removed by adsorbing the iodinated cAMP analog onto a C18 Sep-pak cartridge (Waters, Milford, MA).

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After washing with dH,0, the iodinated cAMP analog was eluted from the Sep-pak Cartridge with 40% acetonitrille (ACN) and 0.1% trifluoroacetic acid (TFA). The iodinated cAMP analog was lyophilized, reconstituted in 1 ml 0.1% 5 TFA, and injected into a C18 reverse phase HPLC column (Waters). The column was equilibrated with 10% ACN in 0.1% TFA, and eluted with gradient of 10-30% ACN in 0.1% This allows separation of the mono-iodinated cAMP analog from the non-iodinated cAMP analog. The tracer is 10 stable for up to 4 months when stored at -20° C. standard used for the assay, adenosine 3':5'-cyclic monophosphate, was purchased from Sigma. Samples (1-10 μ l of HCl extracts) or standards (0.04-100 fmol/tube) were diluted in 50 mM Na-acetate (pH 5.5), and acetylated 15 with 10 μ l of mixture of triethylamine and acetic anhydride (2:1 vol:vol). After acetylation, cAMP antiserum (100 μ l) was added from a stock solution (1:4000) made in PBS (pH 7.4), 5 mM EDTA and 1% normal rabbit serum. The tracer was diluted in PBS (pH 7.4) 20 with 0.1% BSA, and added (20,000 cpm/tube). The assay was incubated at 4° C overnight. The bound tracer was precipitated by adding 100 μ l of goat anti-rabbit antiserum (1:20 in PBS) and 1 ml of 7% polyethyleneglycol (MW 5000-6000), centrifuging at 2000 rpm for 30 min. at The supernatant was removed and the bound radioactivity was counted in a γ -counter (Micromedic). Standard curves were calculated using the four-parameter RIA program supplied by Micromedic. Typically, the assay sensitivity is 0.1 fmol/ tube, and the standard 30 concentration that displaces 50% of tracer is 5

In an alternative method for assaying cAMP accumulation, COS cells transfected with PTH/PTHrP receptor cDNA are harvested with a plastic policeman into a solution containing 10 mM Tris-HCl (pH 7.5), 0.2 mM

fmol/tube.

MgCl, 0.5 mM ethyleneglycolbis(β -amino ethyl ether) N,N'tetra-acetic acid (EGTA) (Sigma) and 1 mM dithiothreitol (Sigma). Cells are homogenated by 20 strokes of tightlyfitting Dounce homogenizer, and centrifuged at 13,000 x q 5 for 15 min at 4°C (Eppendorf, type 5412, Brinkmann Instruments, Inc., Westburg, NY). The pellet containing the plasma membranes is resuspended in the same buffer by several strokes with a Dounce homogenizer, and further diluted with the same buffer to a protein concentration 10 of approximately 1.2 mg/ml, as determined by the method of Lowry et al. (Lowry et al., J. Biol. Chem 193: 265, 1951). Approximately 30 μ g (25 μ l) membrane are incubated with varying concentrations of hormone or vehicle alone for 10 min at 37° C (final volume, 100 μ l) 15 in 50 mM Tris-HC1 (pH 7.5), 0.8 mM ATP, 4 \times 10⁶ cpm [α -32P] ATP (New England Nuclear, Boston, MA), 9 mM theophylline, 4.2 mM MgCl2, 26 mM KCl, 0.12% BSA, and an ATP-regenerating system containing 5 mM creatine phosphate (Schwartz/Mann Division, Becton-Dickenson & 20 Co., Orangeburg, NY) and 0.1 mg/ml creatine phosphokinase (Shwartz/Mann). Incubations are initiated by addition of the membrane suspension and terminated by addition of 100 μ l of a solution containing 20 mM cAMP, approximately 50,000 cpm [3H]cAMP, and 80 mM ATP. The reaction mixture 25 is boiled, and the [32P]cAMP generated is purified by sequential chromatography on ion-exchange columns (Dowex 50 W-X4, Biorad Lab, Richmond, CA) and alumina (Sigma). The [32 P]cAMP may be counted in a β -scintillation counter (Packard Instrument Co.), with correction for recovery of 30 $[^3H]$ CAMP.

Determination of intracellular free calcium

Measurements of intracellular calcium levels in cells transfected with PTH/PTHrP receptor cDNAs were performed using Fura-2 AM (acetomethoxy ester of Fura-2,

Molecular Probes Inc., Eugene, OR) loaded cells. Details of the methodology are:

Coverslips plated with COS cells were incubated in Fura-2 AM loading buffer containing, in mM: HEPES (N-5 [2-Hydroxyethyl]piperazine-N'-[2-ethanesulfonic acid]), 20; $CaCl_2$, 1; KCl 5; NaCl, 145; $MgSO_4$, 0.5; $NaHCO_3$, 25; K_2HPO_4 , 1.4; glucose, 10; and Fura-2 AM 91-(2-5'carboxyoxazol-2'-yl)-6-aminobenzofuran-5oxy-(2'-amino-5'-methylphenoxy) ethane-N, N, N', N'-tetraaecetic acid 10 acetomethoxy ester), 0.5; at 37°C at pH7.4, aerated with 95% air and 5% CO, for 45 minutes. Cells loaded with Fura-2 AM were then washed with a modified Krebs-Heinseleit (KH) buffer containing, in mM: CaCl2, 1; KCl, 5; NaCl, 145; MgSO4, 0.5; Na, HPO4, 1; 15 glucose, 5; pH7.4. To check that cleavage of the ester occurred, the excitation spectra after different times of Fura-2 AM incubation were measured. At 5 min. after the start of incubation, the excitation spectrum peaked at approximately 360 nm, reflecting incomplete hydrolysis of 20 Fura-2 AM, whereas beyond 30 min. the excitation spectrum peaked at 345 nM, characteristic of Fura-2.

To measure fluorescence of individual cells, the cover slips were placed in a microscope tissue chamber (Biophysica Technologies, Inc., MD). The chamber 25 consisted of a shallow, sloped compartment made of Teflon with a silicone rubber seal. The cover slips served as the bottom of the chamber. A heater/cooler ring was encased in the silicone rubber which sealed the coverslip in place. Temperatures were varied between 22°C and 37°C by applying 0-7.4 V to the heater. If the temperature is not specifically stated, the experiment was performed at 37°C. The chamber was mounted on the stage of an inverted microscope (Zeiss IM-35, Thornwood, NY). Fura-2 fluorescence was excited with a 75 watt Xenon arc lamp

35 placed at the focal point of a condenser (Photon

Technologies International (PTI) Inc., NJ). Grating monochromators, alternated by a rotating chopper in which mirror vanes alternate with transmitting sectors, were used for selecting wavelengths. The monochromator outputs were combined to form a common optical path which exited the source housing through an adjustable iris. The light then passed through quartz lenses and a dichroic mirror through a 100x Nikon Fluor objective. A photon-counting PMT device detection was used to measure the light output. Data analysis was performed using PTI software run on an IBM-compatible AT/286 computer using the MS-DOS operating system. Data was retained and

manipulated in a packed binary format. Intracellular calcium concentrations were 15 calculated according to the formula: [Ca²⁺]i=Kd(R-Rmin)/(Rmax-R)B, where R is the ratio of fluorescence of the cell at 340 and 380 nm; Rmax and Rmin represent the ratios of Fura-2 fluorescence intensity at 340 and 380 nm excitation wavelengths in the presence of a saturating 20 amount of calcium and effectively zero calcium, respectively; B is the ratio of fluorescence of Fura-2 at 380 nm in zero calcium to that in saturating amounts of calcium; and $K_{\underline{d}}$ is the dissociation constant of Fura-2 for calcium. To determine Rmax, at the end of an 25 experiment ionomycin was added to the Fura-2 AM loaded cells to equilibrate Ca2+ between the extracellular (1mM) and intracellular environments. To calculate Rmin, 1mM EGTA was then added to the bathing solution. Different

dissociation constants were used at the different temperatures: 224 nM at 34-37°C and 135 nM at 24-27°C.

Determination of inositol phosphate

The level of inositol phosphate metabolism was determined in COS cells transfected with PTH/PTHrP

receptors using previously published methods (Bonventre, et al., J. Biol. Chem. 265: 4934, 1990).

RESULTS

Molecular characterization

Two independent clones (OK-H and OK-O), both of which were isolated from the OK cell cDNA library, had lengths of approximately 2 kilobases. The determined nucleotide sequence and predicted amino acid sequence of these clones are shown in Figs. 1 (SEQ ID NO.:1) and 2 (SEQ ID NO.:2) respectively. The R15B clone isolated from the ROS cell cDNA library had a length of approximately 4 kilobases. The determined nucleotide sequence and predicted amino acid sequence of the rat bone PTH/PTHrP receptor is depicted in Fig. 3 (SEQ ID NO.:3).

The three cDNA clones appear to be full-length by the criteria of having codons encoding methionine residues that are predicted to be the likely candidates as initiator methionines. These methionine codons are followed by amino acid sequences (deduced from the DNA) with properties suggesting that they are "signal-peptide" sequences. All three receptor cDNAs have stop codons at locations that permit these receptors to "fit" a putative seven-membrane spanning model, a model typical for G-protein-linked receptors. Most importantly, all three cloned receptors bind ligands and, when activated, are capable of activating intracellular effectors. These properties suggest that all three of the isolated clones encode full-length cDNAs.

Fig. 4 demonstrates the high degree of homology between the amino acid sequences encoded by the cDNAs from OK-O and ROS 15B. There is an overall 87% homology and a 77.8% amino acid identity between these two receptors. This high level of identity over long

stretches of amnio acids demonstrates that the amino acid sequence of the PTH receptor is evolutionarily conserved to a high degree. This allows the data from both OK-O and R15B to be extrapolated to other species, including 5 human.

Fig. 5 shows the deduced amino acid sequences of all three cloned cDNAs lined up according to sequence homology. The OK-H sequence is identical to OK-O except in the C-terminus tail, where the OK-O sequence totals 10 585 amino acids whereas the OK-H sequence stops at 515 amino acids. This difference is attributable to a single nucleotide (G) deleted in the OK-H sequence compared to the OK-O sequence, causing a frame shift and early stop codon in the former. It is not known whether OK-O and 15 OK-H represent the products of two separate genes or of a laboratory artifact.

Some G-protein-coupled receptors are encoded by intronless genes (Kobilka et al., Nature 329:75, 1987); Kobilka et al., J. Biol. Chem. 262:7321, 1987; Heckert et 20 al., Mol. Endocrinol. 6:70, 1992; Kobilka et al., Science 238:650, 1987; Bonner et al., Science 237:527, 1987; Sunahara et al., Nature 347:80, 1990). To isolate a human PTH/PTHrP receptor cDNA, both a human cDNA library and a human genomic library were screened with a probe 25 (BamHI/NotI) representing most of the coding region of the rat bone PTH/PTHrP receptor (Fig. 3). Screening the human kidney cDNA library led to the isolation of the clone HK-1 (Fig. 6) [SEQ ID NO.: 6]. Since one of the two EcoRI cloning sites of lambda GT10 proved to be 30 eliminated as a result of the library construction, the HindIII/EcoRI phage fragment containing the cDNA insert and ~250 bp of the 37 kb (left) lambda arm was subcloned into the corresponding restriction sites in pcDNAI. DNA sequencing revealed that the cloned cDNA contained ~1000 35 bp of the 3' coding region and ~200 bp of the 3' noncoding region including an A-rich 3' end. The coding region 5' to the XhoI site was subsequently used to rescreen the library and led to the isolation of the clone HK-2 which, after subcloning into pcDNAI, proved to contain ~1400 bp of the coding region. For the third screening of the library, the PvuII/PstI fragment of HK-2 was used; the isolated clone HK-3 proved to be identical to HK-2.

The genomic library screening (~10⁶ pfu) resulted in the isolation of four independent clones. Comparison of Southern blot analyses of restriction enzyme digests of these clones with that of normal genomic DNA, revealed that one 15 kb genomic clone, HPG1 (also referred to as HG4A), contained a SstI/SstI fragment that had the same size as one hybridizing DNA species from normal human genomic DNA digested with SstI (see heles).

genomic DNA digested with SstI (see below). The hybridizing 2.3 kb SstI/SstI DNA fragment and an ~8 kb XhoI fragment which comprised the SstI/SstI fragment were both subcloned into pcDNAI. Further Southern blot

analysis of the SstI/SstI DNA fragment revealed that an ~1000 bp BamHI/SstI fragment encoded a portion of the human PTH/PTHrP receptor which later proved to represent the exon encoding the putative signal peptide and the 5' non-translated region which is interrupted by an ~1000 bp intron (Fig. 7).

To isolate the remaining ~450 nucleotides of the coding region, poly (A)+ RNA from human kidney was reverse transcribed after priming with H12 (Fig. 7). After single strand synthesis, two independent PCRs were performed using two different forward primers: i) a degenerate primer RK- 1 based on the 5' coding end of the two previously cloned PTH/PTHrP receptors, OK-O and R15B; and ii) primer RK-2 based on the 5' non-coding region of HPG1. H-26 was used as the reverse primer for both

35 reactions. Southern blot and restriction map analyses

confirmed the expected size of the amplified DNA encoding the human PTH/PTHrP receptor. The blunt-ended PCR products encoding the 5' end of the human PTH/PTHrP were cloned into pcDNAI using the dephosphorylated EcoRV sites. Sequence analysis of each PCR clone confirmed their 5' nucleotide difference due to the difference in forward primer sequence, but revealed otherwise identical sequences. Nucleotide sequencing of both strands of the human PTH/PTHrP receptor cDNA revealed an open reading frame encoding a 593-amino acid protein (Fig. 6, SEQ ID NO.:4).

The full-length human kidney PTH/PTHrP receptor cDNA, HKrk, was constructed using the BamHI/PvuII fragment of PCR clone #2 and HK-2. Using the full-

- length cDNA encoding the human PTH/PTHrP receptor,
 Northern blot analysis of total RNA (~10 µg/lane) from
 human kidney and SaOS-2 cells revealed one major
 hybridizing DNA species of ~2.5 kb (Fig. 19). The XhoI
 digest of normal human genomic DNA, when probed with the
- same full-length cDNA (Fig. 20), revealed one major hybridizing species of about 5.5 kb, and two DNA species of 4 and 8 kb which weakly hybridized. These date suggest that the human PTH/PTHrP receptor is the product of a single gene. This full-length clone was then
- 25 transiently expressed in COS-7 cells for functional and biological characterization by the methods cited above.

Comparison of the human receptor with the opposum kidney PTH/PTHrP receptor and the rate bone PTH/PTHrP receptor, revealed 81% and 91% amino acid sequence

identity, respectively, and consequently a very similar hydrophobicity plot (Fig. 8). All extracellular cysteines including the two cysteine residues in the presumed signal peptide are conserved, as are all potential, extracellular

N-glycosylation sites. A number of the amino acids which were not identical between the human kidney and rat bone PTH/PTHr receptors were found to be conserved between the human and the opposum receptors. These conserved amino acids include an Arg to Leu at 51, an Arg to Trp at 58, an Arg to His at 262, an Asp to His at 358, an Ile to Thr at 422, and a Thr to Leu at 427.

Biological Characterization

Functional characterization of the biological
properties of the opossum and rat PTH/PTHrP receptors was
performed in transiently transfected COS cells by a
radioreceptor assay technique using both ¹²⁵I-PTHrP and
¹²⁵I-NlePTH as radioligands, and by bioassays that measure
ligand-stimulated cAMP accumulation, increase in

15 intracellular free calcium, and stimulation of inositol phosphate metabolism, by the methods cited above.

Fig. 9 demonstrates that COS cells expressing OK-H bind ¹²⁵I-PTHrP. These data also demonstrate that binding of PTHrP is inhibited when intact PTH (1-34) or PTH anlogues which are shortened at their amino terminus (i.e. the 3-34 and 7-34 analogues, which contain Nle substitutions for methionine at positions 8 and 18 and a tyrosine substitution for phenylalanine at position 34) are used as competitors for binding. Similarly, binding of ¹²⁵I-NlePTH to COS cells expressing OK-H was inhibited when PTHrP or PTHrP fragments were used as competitors. These data indicate that PTH and PTHrP both bind to the receptor encoded by OK-H.

Fig. 10 demonstrates that COS cells expressing OK-30 H increase their concentration of intracellular free calcium when exposed to NlePTH, but to a smaller extent (mean =

39 nm), or not at all, when compared to COS cells expressing OK-O or R15B receptors (Fig. 12 and Fig. 14)

35 and stimulated with NlePTH. Unlike COS cells expressing

OK-O or R15B, COS cells expressing OK-H do not show a detectable increase in metabolism of inositol phosphate when stimulated with NlePTH (Fig. 15).

Fig. 11 demonstrates that COS cells expressing OK5 O bind ¹²⁵I-PTHrP. These data also demonstrate that
binding of PTHrP is inhibited when intact PTH (1-34) or
PTH analogues which are shortened at their amino terminus
(i.e. the 3-34 and 7-34 analogues, which contain Nle
substitutions for methionine at positions 8 and 18 and a
10 tyrosine substitution for phenylalanine at position 34)
are used as competitors for binding. Similarly, binding
of ¹²⁵I-NlePTH to COS cells expressing OK-H was inhibited
when PTHrP or PTHrP fragments were used as competitors.
These data indicate that PTH and PTHrP both bind to the
15 receptor encoded by OK-O.

Fig. 12 demonstrates that COS cells expressing OK-O increase their concentration of intracellular free calcium and their rate of inositol phosphate metabolism after stimulation with NlePTH and PTHrP (Fig. 15).

Fig. 13 demonstrates that COS cells expressing R15B bind ¹²⁵I-PTHrP. These data also demonstrate that binding of PTHrP is inhibited when intact PTH (1-34) or PTH anlogues which are shortened at their amino terminus (i.e. the 3-34 and 7-34 analogues, which contain Nle substitutions for methionine at positions 8 and 18 and a tyrosine substitution for phenylalanine at position 34) are used as competitors for binding. Similarly, binding of ¹²⁵I-NlePTH to COS cells expressing OK-H was inhibited

when PTHrP or PTHrP fragments were used as competitors.

These data indicate that PTH and PTHrP both bind to the receptor encoded by R15B.

Fig. 14 demonstrates that COS cells expressing R15B increase their concentration of intracellular calcium to an extent similar to stimulated COS cells expressing OK-O.

Fig. 15 demonstrates that COS cells expressing R15B or OK-O increase their rate of phosphatidyl inositol hydrolysis, as evidenced by the rapid increase in inositol trisphosphate (IP3) and inositol bisphosphate 5 (IP2) accumulation after stimulation of the cells with NlePTH or PTHrP. Conversely, COS cells expressing OK-H did not show any detectable increase in inositol trisphosphate and inositol bisphosphate accumulation after stimulation with NlePTH or PTHrP. These data 10 suggest that the PTH receptor encoded by R15B and OK-O is coupled to phospholipase C, presumably through Gp. Since the only difference between OK-O and OK-H is in the cytoplasmic C-terminal tail, these data strongly suggest that the C-terminus of the PTH receptor encoded by OK-O 15 and R15B is involved in the activation of phospholipase c.

Fig. 16 demonstrates that COS cells expressing R15B and OK-H increase cAMP accumulation after stimulation with NlePTH. Similar results were obtained 20 in COS cells expressing OK-O. No cAMP stimulation was detected in COS cells transfected with the cDM8 vector These data suggest that PTH receptor coupling to adenylate cyclase does not require the full length Cterminal cytoplasmic tail of the receptor.

These data demonstrate that all three PTH/PTHrP receptors cloned from both OK and ROS cell cDNA libraries bind the amino-terminal ligands of both peptides equivalently. Activation of all these receptors by ligand stimulates adenylate cyclase (as measured by 30 increased intracellular cAMP), presumably through activation of one class of guanine nucleotide binding proteins (G-proteins). G-proteins have a trimeric peptide structure in which one of the subunits, alpha, is distinct, and the other two, beta and gamma, are 35 identical or highly homologous. One of these G-proteins

25

2.000

 (G_s) contains G-alpha-"stimulatory" (G-alpha-s) which is involved in the activation of adenylate cyclase.

Binding of ligand to OK-O and R15B, but not to OK-H, also increases intracellular free calcium and

5 stimulates metabolism of inositol phosphate. These properties strongly suggest that activation of both OK-O and R15B receptors by ligand results in stimulation of a second intracellular effector, phospholipase C. The coupling mechanism between these activated receptors and phospholipase C is likely to be a G-protein which is distinct from G_s. In contrast, the properties of the activated OK-H receptor which is truncated at the carboxy terminus, suggest that it may not activate phospholipase C, or that it activates phospholipase C inefficiently.

The biochemical role of the carboxy-terminal tail 15 of the PTH/PTHrP receptor was further investigated by the construction of a carboxy-terminally-truncated rat receptor, R480, by standard PCR technology using R15B as a template and an upstream primer containing a stop codon 20 inserted at position 481. Briefly, the upstream primer was a synthetic oligonucleotide based on nucleotides 1494-1513 of the rat cDNA sequence (see Fig. 3; SEQ ID NO.: 3) to which a stop codon and an XbaI cloning site were added. Thirty PCR cycles were carried out, each 25 cycle consisting of 1 min at 92°C for denaturation, 1 min at 60°C for annealing, and 1 min at 72°C for extension. The product was cut with NsiI and XbaI and purified by gel electrophoresis. R15B was sequentially digested with XbaI and NsiI, and the purified PCR product was then 30 ligated into the XbaI-NsiI cut R15B vector. resulting plasmid, R480, was amplified in bacteria and sequenced.

R480 encodes 480 amino acids that are identical to those in the 591 amino acids receptor. This truncated 35 cDNA was expressed in COS-7 cells (transient expression)

and in CHO cells (stable expression). Both COS-7 and CHO cells expressing the truncated receptor, R480, and the wild type receptor, RB, bind PTH(1-34) with equivalent affinities. When activated, R480 stimulates cAMP 5 accumulation in COS7 and CHO cells as efficiently as does the wild type receptor. In contrast to the wild type receptor, R480 did not mediate any increase in [Ca2+]i when stimulated by PTH in either the COS-7 cells or the CHO cells. These data indicate that the molecular 10 requirements for activation of phospholipase C and adenylate cyclase by PTH/PTHrP receptor are distinct from each other, and point to a major role of the carboxyterminal tail of the PTH/PTHrP receptor in coupling to phospholipase C but not to adenylate cyclase. Of course, 15 it is also possible that activated PTH/PTHrP receptors may activate additional G-proteins and/or intracelluar effector molecules.

Analysis of COS-7 cells transfected with the cloned human PTH/PTHrP receptor demonstrated that 20 radiolabelled PTH(1-34) and PTHrP(1-36) (~200,000 cpm) bound to the expressed receptors with similar efficiency (specific binding: $10.1 \pm 3.7\%$ and $7.6\pm6.0\%$, respectively) to that observed for COS-7 cells expressing R15B (specific binding: 8.1+3.5% and 7.1+4.1%, 25 respectively). The expressed human PTH/PTHrP receptors bound PTH(1-34) with 2-fold higher apparent Kd than did the rat bone PTH/PTHrP receptor: ~5 nM versus ~10 nM (Fig. 17). However, despite their high degree of amino acid homology, the two receptors showed significant 30 differences in affinity for PTH(3-34) and PTH(7-34). PTHrP(1-36) displayed a 2- to 4-fold lower affinity for the human PTH/PTHrP receptor than for the rat receptor (~35 nM for HKrk versus ~10 nM for R15B) which appeared more pronounced when PTHrP(1-36) was used as radioligand. 35 The affinities for PTH(3-34) and PTH(7-34) were 7- and

35-fold higher with the expressed HKrK than with R15B (~7 nM versus ~45 nM for PTH(3-34), respectively; ~60 nM versus ~2000 nM for PTH(7-34), respectively). In COS-7 cells expressing either receptor, both PTH(1-34) and PTHrP(1-36) stimulated the increase in intracellular free calcium and cAMP accumulation to the same extent (Fig. 18).

Relationship of PTH/PTHrP receptors

- The amino acid sequence of the human PTH/PTHrP

 10 receptor displays a very high degree of conservation compared to the bone PTH/PTHrP receptor from rat, a eutherian mammal, while its sequence identity with the PTH/PTHrP receptor with the opossum, a marsupial mammal, is less marked. Like the opossum kidney and the rat bone
- 15 receptor, the human kidney receptor induces an increase in both intra-cellular cAMP and intracellular free calcium when challenged with either PTH or PTHrp.

 Despite the high degree of homology between the human PTH/PTHrP receptor and the opossum and rat homologs, the
- transiently expressed human receptor has some functional characteristics that are distinct from those of the rat bone receptor. These include a slightly higher affinity for PTH(1-34) and a significantly descreased affinity for PTHrP(1-36). Higher affinities were observed for PTH(3-
- 25 34) and in particular for PTH(7-34), the affinity of which for the human receptor was about 35-fold higher in comparison to the rat bone receptor. These findings may have significant implications for the future development of PTH/PTHrP analogues, since they predict that species-
- 30 specific tissues would be the appropriate tissues for testing the potency of antagonists (and agonists) in vitro.

Relationship of PTH/PTHrP receptors to other receptors

The biochemical properties of PTH and PTHrP

35 receptors suggest that they are members of the class of

membrane receptor molecules known as G-protein-linked membrane receptors. The structural features of well-characterized G-protein receptors indicate that they all have at least seven regions of several consecutive hydrophobic amino acids, each of which regions is of sufficient length to span the plasma membrane.

One subfamily of G-protein-linked membrane receptors, termed the glycopeptide receptor subfamily, includes receptors that bind and are activated by 10 glycopeptide hormones (thyroid-stimulating hormone, luteinizing hormone, follicle-stimulating hormone, and chorionic gonadotropin). All of these receptors are characterized by (1) extensive putative amino-terminal extracellular domains (greater than 300 amino acids) that 15 are thought to contain some or all of the ligand-binding domains, and (2) considerable amino-acid homology, particularly in the seven putative transmembrane domains. A second subfamily, termed the adrenergic/muscarinic subfamily, includes receptors that are activated by small 20 ligands, such as the catecholoamines, neuromuscular transmitters, and retinol. These receptors are all characterized by relatively short (25-75 amino acids) putative amino-terminal extracellular domains, as well as considerable amino acid homology, particularly in the 25 seven putative transmembrane domains. Activation of these receptors by their ligands appears to involve at least several of the multiple transmembrane domains, and does not appear to involve the amino-terminal portion of the receptors.

30 Several structural characteristics which can be deduced from the predicted amino acid sequence of the rat PTH/PTHrP receptor (Fig. 3) indicate that the PTH/PTHrP is a G-protein-linked receptor. The amino terminus shows characteristic features of a signal peptide, including a hydrophobic domain and the presence of three consecutive

leucine residues. This amino acid stretch of 20-28 amino acids may serve as a leader sequence, similar to the amino terminus preceding the extracellular domains of other glycoprotein receptors. There is also a cluster of seven hydrophobic segments which represent putative membrane-spanning domains (Fig. 19).

The predicted amino acid sequences of the opossum kidney, rat bone and human kidney PTH/PTHrp receptors indicate that they do not fit comfortably into either of 10 these G-protein linked receptor subfamilies. Overall homology of the rat and human PTH/PTHrP receptors with the glycopeptide receptor and adrenergic/muscarinic subfamilies is approximately 10 to 20%, with a somewhat higher degree of homology within the transmembrane 15 domains. The latter is to be expected because of the limited menu of hydrophobic amino acids that could occur in those regions. Twenty percent homology is far less than that found among the receptors generally accepted to be members of each of these subfamilies. Additionally, 20 there are no portions of these sequences that have what could be characterized as intense homology (i.e., exactly matching amino acid sequences), even over limited regions.

Recent comparison with the newly characterized

25 secretin and calcitonin receptors (Ishihara et al., EMBO

J 10:1635, 1991; Lin et al., Science 254:1022, 1991) has
revealed between 30 and 40% identity between these
receptors and the PTH/PTHrP receptor. Although the
PTH/PTHrP receptor is more than 100 amino acids longer

30 than the calcitonin receptor, there is an ~32% identify
between the amino acid sequences of the opossum kidney
PTH/PTHrP receptor (SEQ ID NO NO.:2) and porcine kidney
calcitonin receptor (GenBank accession no. M74420). A
stretch of 17 out of 18 amino acids in the putative

35 transmembrane domain VII are identical. Also, two out of

four N-linked glycoslyation sites and the position of seven out of eight potentially extracellular cysteines are conserved. Major differences between the two receptors appear to lie in their NH,-terminal and COOH-5 terminal domains. Comparison of amino acid sequences of the rat secretin receptor (GenBank accession no. X59132) and the human PTH/PTHrP receptor indicates that there is a 43% identity between these two receptors, with a stretch of 21 out of 25 amino acids of the putative 10 transmembrane domain VII being identical. The similarity between the PTH/PTHrP, calcitonin and secretin receptors suggests that they represent a new family of seven transmembrane-spanning G protein-coupled receptors that activate adenylate cyclase. Given the amino acid 15 sequences of these receptors, those skilled in the art would be able to compare these sequences for regions of identity which would be useful in the design of nucleic acid probes which could then be used for the identification and isolation of other receptors which 20 would belong to this family.

Deposit of Clones

Under the terms of the Budapest Treaty on the
International Recognition of the Deposit of
Microorganisms for the Purpose of Patent Procedure, the
25 cDNA expression plasmids R15B, OK-O, and OK-H; the phage
HPG1; and a plasmid (termed 8A6) containing part of the
human clone have been deposited with the American Type
Culture Collection (ATCC), where they bear the respective
accession numbers ATCC No. 68571, 68572, 68573, 40998 and
30 68570. Applicants' assignee, The General Hospital
Corporation, represents that the ATCC is a depository
affording permanence of the deposits and ready
accessibility thereto by the public if a patent is
granted. All restrictions on the availability to the

public of the material so deposited will be irrevocably removed upon the granting of a patent. The material will be available during the pendency of the patent application to one determined by the Commissioner to be 5 entitled thereto under 37 CFR 1.14 and 35 U.S.C. 122. The deposited material will be maintained with all the care necessary to keep it viable and uncontaminated for a period of at least five years after the most recent request for the furnishing of a sample of the deposited 10 plasmid, and in any case, for a period of at least thirty (30) years after the date of deposit or for the enforceable life of the patent, whichever period is longer. Applicants' assignee acknowledges its responsibility to replace the deposits should the 15 depository be unable to furnish a sample when requested due to the condition of the deposit. **POLYPEPTIDES**

Polypeptides according to the invention include the opossum and rat and human parathyroid hormone

receptors as shown in Figs. 1-3 and 6, respectively, and any other naturally-occurring receptor which can be produced by methods analogous to those used to clone and express these receptors, or by methods utilizing as a probe all or part of one of the sequences described herein. In addition, any analog or fragment of a PTH receptor capable of binding to a parathyroid hormone or a parathyroid hormone-related protein is within the invention.

Specific receptor analogs of interest include

30 full-length or partial receptor proteins having an amino acid sequence which differs only by conservative amino acid substitutions: for example, substitution of one amino acid for another of the same class (e.g., valine for glycine; arginine for lysine, etc.), or by one or

35 more non-conservative amino-acid substitutions,

deletions, or insertions located at positions which do not destroy the receptor's ability to bind to parathyroid hormone or parathyroid hormone-related protein.

Specific receptor fragments of particular interest include, but are not limited to, portions of the receptor deduced to be extracellular from the primary amino acid sequence, using a hydrophobicity/hydrophilicity calculation such as the Chou-Fasman method (see, e.g., Chou and Fasman, Ann. Rev. Biochem. 47:251, 1978).

- 10 Hydrophilic domains, particularly ones surrounded by hydrophobic stretches (e.g., transmembrane domains) of at least 10 amino acids, present themselves as strong candidates for extracellular domains. Fig. 21 illustrates a predicted arrangement of extracellular,
- 15 intracellular, and transmembrane domains of one PTH receptor.

Examples of specific PTH receptor fragments include those with the following amino acid sequences (shown as standard single-letter symbols), derived from the deduced amino acid sequence of the R15B clone:

Extracellular domains:

RP-1: TNETREREVFDRLGMIYTVG (SEQ ID NO.: 5)

RP-2: VLYSGFTLDEAERLTEEEL (SEQ ID NO.: 6)

RP-3: VTFFLYFLATNYYWILVEG (SEQ ID NO.: 7)

25 RP-4: Y-RATLANTGCWDLSSGHKKWIIQVP (SEQ ID NO.: 8)

RP-5: PYTEVSGTLWQIQMHYEM (SEQ ID NO.: 9)

RP-6: DDVFTKEEQIFLLHRAQA (SEQ ID NO.: 10)

Intracellular domains:

RPi-7: FRRLHCTRNY (SEQ ID NO.: 11)

30 RPi-8: EKKYLWGFTL (SEQ ID NO.: 12)

RPi-9: VLATKLRETNAGRCDTRQQYRKLLK (SEQ ID NO.: 13)
These fragments were synthesized and purified by HPLC according to the method of Keutmann et al.,
(Endocrinology 117: 1230, 1984).

EXPRESSION OF POLYPEPTIDES

Polypeptides according to the invention may be produced by expression from a recombinant nucleic acid having a sequence encoding part or all of a cell receptor 5 of the invention, using any appropriate expression system: e.g., transformation of a suitable host cell (either prokaryotic or eukaryotic) with the recombinant nucleic acid in a suitable expression vehicle (e.g., pcDNAI). The precise host cell used is not critical to 10 the invention; however, in the case wherein the polypeptides of the invention include all or part of the PTH/PTHrP receptor, the following host cells are preferred: COS cells, LLC-PK1 cells, OK cells, AtT20 cells, and CHO cells. The method of transfection and the 15 choice of expression vehicle will depend on the host system selected. Mammalian cell transfection methods are described, e.g., in Ausubel et al. (Current Protocols in Molecular Biology, John Wiley & Sons, New York, 1989); expression vehicles may be chosen from those discussed, 20 e.g., in Cloning Vectors: A Laboratory Manual (P.H. Pouwels et al., 1985, Supp. 1987). Stably transfected cells are produced via integration of receptor DNA into the host cell chromosomes. Suitable DNAs are inserted into pcDNA, pcDNAI-Neo, or another suitable plasmid, and 25 then cells are transfected with this plasmid with or without cotransfection with psV-2-Neo, or psV-2-DHFR by standard electroporation, calcium phosphate, and/or DEAE/Dextran techniques. Selection of transfected cells is performed using progressively increasing levels of 30 G418 (Geneticin, GIBCO), and if necessary, methotrexate.

DNA sequences encoding the polypeptides of the invention can also be expressed in a prokaryotic host cell. DNA encoding a cell receptor or receptor fragment is carried on a vector operably linked to control signals capable of effecting expression in the prokaryotic host.

If desired, the coding sequence may contain, at its 5' end, a sequence encoding any of the known signal sequences capable of effecting secretion of the expressed protein into the periplasmic space of the host cell,

- 5 thereby facilitating recovery of the protein and subsequent purification. Prokaryotes most frequently used are various strains of <u>E. coli</u>; however, other microbial strains may also be used. Plasmid vectors are used which contain replication origins, selectable
- 10 markers, and control sequences derived from a species compatible with the microbial host. For example, <u>E. coli</u> may be transformed using derivatives of pBR322, a plasmid constructed by Bolivar et al. (Gene 2: 95, 1977) using fragments derived from three naturally-occurring
- plasmids, two isolated from species of <u>Salmonella</u>, and one isolated from <u>E. coli</u>. pBR322 contains genes from ampicillin and tetracycline resistance, and thus provides multiple selectable markers which can be either retained or destroyed in constructing the desired expression
- vector. Commonly used prokaryotic control sequences
 (also referred to as "regulatory elements") are defined
 herein to include promoters for transcription initiation,
 optionally with an operator, along with ribosome binding
 site sequences. Promoters commonly used to direct
- protein expression include the beta-lactamase (penicillinase), the lactose (lac) (Chang et al., Nature 198: 1056, 1977) and the tryptophan (Trp) promoter systems (Goeddel et al., Nucl. Acids Res. 8: 4057, 1980) as well as the lambda-derived P_L promoter and N-gene
- 30 ribosome binding site (Simatake et al., Nature 292:128, 1981).

The nature of the cell receptor proteins of the invention is such that, upon expression within a cell, it is moved to the cellular membrane and partially through the membrane, so that part of it remains embedded in the

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membrane, part extends outside the cell, and part remains within the cell. Transformed cells bearing such embedded cell receptors may themselves be employed in the methods of the invention, or the receptor protein may be extracted from the membranes and purified.

Expression of peptide fragments lacking the hydrophobic portions of the protein responsible for anchoring the intact protein in the cellular membrane would not be expected to become embedded in the membrane; whether they remain within the cell or are secreted into the extracellular medium depends upon whether or not a mechanism promoting secretion (e.g., a signal peptide) is included. If secreted, the polypeptide of the invention can be harvested from the medium; if not, the cells must be broken open and the desired polypeptide isolated from the entire contents of the cells. Specific examples of polypeptides which might be expressed include, without limitation:

- Amino-terminal portion comprising amino acids
 1-192, including the putative leader sequence, of the rat bone PTH/PTHrP receptor as shown in Fig. 3.
 - 2) Amino-terminal portion comprising amino acids 27-192, excluding the putative leader sequence, of the rat bone PTH/PTHrP receptor as shown in Fig. 3.
- 25 3) The full-length PTH/PTHrP receptor from rat bone, as shown in Fig 3.
 - 4) RP-1 (as described above).
 - 5) RP-2 (as described above).

The polypeptide of the invention can be readily
purified using affinity chromatography. Antibodies to
these polypeptides, or the receptor specific ligands,
(e.g., the hormones PTH and PTHrP for the PTH/PTHrP
receptor) may be covalently coupled to a solid phase
support such as Sepharose 4 CNBr-activated sepharose
(Pharmacia), and used to separate the polypeptide of the

invention from any contaminating substances. Typically 1 mg of ligand or antibody will be incubated with CNBractivated sepharose at 4°C for 17-20 h (with shaking). The sepharose is rinsed with 1 M Tris HCL (pH8) to block excess active sites. The sepharose-PTH, sepharose-PTHrP, or sepharose-antibody is then incubated with the crude polypeptide in phosphate-buffered saline (pH 7.4) at 4°C for 2 h (with shaking). The sepharose is then typically packed in a column, thoroughly washed with PBS (typically 10 times the column volume), and eluted with dilute HCl in H₂O (pH 1.85). The eluate may then be concentrated by lyophylization and its purity checked, for example, by reverse phase HPLC.

ANTI-CELL RECEPTOR ANTIBODIES

Cell receptor or receptor fragments of the 15 invention may be used to generate antibodies by any conventional method well known to those skilled in the art, including those which generate polyclonal antibodies and those which generate monoclonal antibodies. 20 example, the deduced amino acid sequence of the PTH receptor reveals a protein structure that appears to have several transmembrane (i.e., hydrophobic) domains interspersed with presumably extracellular and intracellular regions (see Fig. 21) analogous to those 25 found in other G protein-linked receptors. This information can be used to guide the selection of regions of the receptor protein which would be likely to be exposed on the cell surface, and thus would be presented to antibodies in vivo. A short peptide representing one 30 or more of such regions may be synthesized (e.g., chemically or by recombinant DNA techniques) and used to immunize an animal (e.g., a rabbit or a mouse) to generate polyclonal or monoclonal antibodies. For example, certain of the peptides of the PTH/PTHrP 35 receptor listed above (RP-1, RP-5 and RP-6) have been

chemically synthesized using standard techniques and used to generate polyclonal antibodies in rabbits by the following procedure:

A preparation of a given peptide emulsified with 5 complete Freund's Adjuvant is injected intradermally into rabbits. Booster injections are emulsified in or complete adjuvant and injected at monthly intervals.

Antibody titer is assessed using either of two methods. First, serial dilutions of the antiserum in 1% normal rabbit serum are incubated with 125 I-labelled PTH/PTHrP receptor fragment by standard methods (e.g., see Segre et al., supra) for 24 h at 4° C. The bound 125 I-PTH/PTHrP receptor fragments are separated from unbound by addition of 100 μ l of second antibody (anti-rabbit IgG, Sigma) diluted 1:20 and 1 ml of 5%

- polyethylene glycol, followed by centrifugation at 2000 rpm for 30 min. at 4° C. The supernatant is removed and the pellet analyzed for radioactivity in a γ -counter. In the second method, cell lines expressing either native
- 20 (e.g., ROS 17/2.8, OK, SaOS-02 cells) or recombinant (COS cells or CHO cells transfected with R15B, OK-O or OK-H) PTH/PTHrP receptors are incubated with serially diluted antibody at 4°C, 20°C or 37°C for
- 1- 4 h. The cells are rinsed with PBS (x3) and incubated for 2 h at 4°C with 125 I-labelled (NEN, Dupont) or FITC-labelled (Sigma) second antibodies. After rinsing (x3 with PBS), the cells were either lysed with 0.1 M NaOH and counted in γ -counter (if 125 I-labelled second antibody was used) or fixed with 1% paraformaldehyde and examined
- 30 by fluorescent microscopy (if FITC-labelled second antibody was used).

Another method for producing antibodies utilizes as antigen the intact cell receptor protein of the invention expressed on the surface of cells (e.g., 35 mammalian cells, such as COS cells, transfected with DNA

encoding the receptor). Such cells are prepared by standard techniques, e.g., by the DEAE-dextran transfection method, using a vector encoding and capable of directing high-level expression of the cell receptor.

5 Such cells may be used to generate polyclonal or monoclonal antibodies. For example, monoclonal antibodies specific for the PTH/PTHrP receptor may be produced by the following procedure:

Intact COS cells expressing high levels of rat 10 recombinant PTH receptors on the cell surface are injected intraperitoneally (IP) into Balb-c mice (Charles River Laboratories, Willmington, MA). The mice are boosted every 4 weeks by IP injection, and are hyperimmunized by an intravenous (IV) booster 3 days 15 before fusion. Spleen cells from the mice are isolated and are fused by standard methods to myeloma cells. Hybridomas are selected in standard hypoxanthine/aminopterin/thymine (HAT) medium, according to standard methods. Hybridomas secreting antibodies 20 which recognize the PTH receptor are initially identified by screening with cell lines which naturally express abundant copies of the PTH-receptor per cell (such as ROS17/2.8 or OK cells), using standard immunological techniques. Those hybridomas which produce antibodies 25 capable of binding to the PTH receptor are cultured and subcloned. Secondary screening with radioreceptor and cAMP stimulation assays can then be performed to further characterize the monoclonal antibodies (see below). SCREENING FOR PTH RECEPTOR ANTAGONISTS AND AGONISTS

The polypeptides and antibodies of the invention and other compounds may be screened for PTH-competition and for antagonistic or agonistic properties using the assays described herein.

In one example, those antibodies that recognize 35 the PTH receptor on the intact cells are screened for

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their ability to compete with PTH or PTHrP for binding to a PTH/PTHrP receptor. Cells expressing PTH receptor on the cell surface are incubated with the ¹²⁵I-PTH analog, ¹²⁵I-NlePTH or ¹²⁵I-PTHrP in the presence or absence of the polyclonal or monoclonal antibody to be tested, for 4 h at 15°C. The antibody used may be from crude antiserum, cell medium, or ascites, or in purified form. After incubation, the cells are rinsed with binding buffer (e.g., physiological saline), lysed, and quantitatively analyzed for radioactivity using a gammacounter. Antibodies that reduce binding of the PTH analog to the PTH receptor are classified as competitive; those which do not are noncompetitive.

Compounds, including antibodies and polypeptides, 15 may be screened for their agonistic or antagonistic properties using the cAMP accumulation, intracellular calcium, and/or inositol phosphate assays described above. Cells expressing PTH receptor on the cell surface are incubated with PTH, PTH-receptor antibody, or a 20 combination of both, for 5 - 60 minutes at 37°C, in the presence of 2 mM IBMX (3-isobutyl-1-methyl-xanthine, Sigma, St. Louis, MO). Cyclic AMP accumulation is measured by specific radio-immunoassay, as described above. A compound that competes with PTH for binding to 25 the PTH receptor, and that inhibits the effect of PTH on cAMP accumulation, is considered a competitive PTH antagonist. Conversely, a compound that does not compete for PTH binding to the PTH receptor, but which still prevents PTH activation of cAMP accumulation (presumably 30 by blocking the receptor activation site) is considered a non-competitive antagonist. A compound that competes with PTH for binding to the PTH receptor, and which stimulates cAMP accumulation in the presence or absence of PTH, is a competitive agonist. A compound that does 35 not compete with PTH for binding to the PTH receptor but

which is still capable of stimulating cAMP accumulation in the presence or absence of PTH, or which stimulates a higher accumulation than that observed by PTH alone, would be considered a non-competitive agonist.

5 USE

The polypeptides, antibodies, and other compounds of the invention are useful for the diagnosis, classification, prognosis, and/or treatment of disorders which may be characterized as related to the interaction 10 between a cell receptor of the invention and its specific ligand. For example, some forms of hypercalcemia and hypocalcemia are related to the interaction between PTH and PTHrP and the PTH/PTHrP receptor(s). Hypercalcemia is an condition in which there is an abnormal elevation 15 in serum calcium level; it is often associated with other diseases, including hyperparathyroidism, osteoporosis, carcinomas of the breast, lung and prostrate, epidermoid cancers of the head and neck of the esophagus, multiple myeloma, and hypernephroma. Hypocalcemia, a condition in 20 which the serum calcium level is abnormally low, may result from a deficiency of effective PTH, e.g., following thyroid surgery.

In a first example, the compounds of the invention are used to manufacture diagnostic agents which are used as diagnostic tools to diagnose hypercalcemia and to distinguish between hypercalcemic conditions, i.e., to differentiate hypercalcemia mediated by PTH or PTHrP (e.g., hyperparathyroidism and humoral hypercalcemia of malignancy), from hypercalcemia associated with diseases which do not involve these factors (e.g., local osteolytic hypercalcemia mediated by the presence of metastatic tumor cells in direct contact with bone, and certain rare types of malignancy-related hypercalcemias mediated by an increase of humoral factors, such as osteoclast activating factor (interleukin), lymphotoxin,

calcitriol, type E prostaglandins, and vitamin D-like sterols).

In one method of diagnosis, serum total and/or ionized calcium levels are measured by standard

5 techniques before and after the administration of the PTH or PTHrP antagonists of the invention. PTH or PTHrP related hypercalcemias would be detectable as a decrease in serum calcium levels following administration of the antagonist of the invention. In contrast, for

10 hypercalcemic conditions mediated by factors other than PTH or PTHrP, the serum calcium levels would remain

unchanged even after administration of the antagonist.

Another diagnostic application of the invention permits measurement of the level of PTH or PTHrP in a 15 biological sample in order to diagnose PTH or PTHrP related tumors, e.g., tumors which are associated with humoral hypercalcemia of malignancy, and for monitoring . the levels of PTH or PTHrP during cancer therapy. method involves assaying binding of the recombinant 20 parathyroid hormone receptor of the invention to PTH or PTHrP present in a tissue sample, using the binding assay described herein. The level of binding may be determined directly (e.g., by using radioactively labelled PTH receptor, and assaying the radioactivity bound to 25 endogenous PTH). Alternatively, binding of PTH receptor to the sample (e.g., a tissue section) may be followed by staining of the tissue sections with an antibody specific for the PTH receptor, using standard immunological techniques (Chin et al., Hybridoma 5:339, 1986).

In a third diagnostic approach, one could stably transfect cell lines (by the methods described in Ausubel et al., <u>Current Protocols in Molecular Biology</u>, Wiley Publishers, New York, 1987) with a PTH receptor gene linked to an appropriate promoter (e.g., the

35 metallothionine promoter). Alternatively, the PTH/PTHrP

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receptor could be expressed from a eukaryotic vector, i.e., pcDNAI, and cotransfected with a mutant DHFR gene that will allow further gene amplification via methotrexate selection (Simonsen et al., Proc. Natl. 5 Acad. Sci., <u>80</u>:2495-2499, 1983). Such high-level expression of the gene produces an immortal cell line which is oversensitive to PTH or PTHrP. Such cells provide a particularly useful tool for detecting serum blood levels of PTH or PTHrP. Such a cell line may be 10 used for diagnosis of conditions involving elevated PTH or PTHrP levels (e.g., those described above) or for conditions involving unusually low levels of PTH or PTHrP (e.g., those described above). Such a cell line is also useful for monitoring the regression or increase of PTH 15 or PTHrP levels during therapy for hypercalcemia or hypocalcemia, respectively.

A patient who is suspected of being hypercalcemic may be treated using the compounds of the invention. Rapid intervention is important because symptoms may 20 appear abruptly and, unless reversed, can be fatal. one application, serum calcium levels are stabilized by an immediate course of treatment which includes antagonists of PTH or PTHrP. Such antagonists include the compounds of the invention which have been determined 25 (by the assays described herein) to interfere with PTH receptor-mediated cell activation. To administer the antagonist, the appropriate antibody or peptide (is used in the manufacture of a medicament, generally by being formulated in an appropriate carrier such as 30 physiological saline, and administered intravenously, at a dosage that provides adequate competition for PTH or PTHrP binding to the PTH receptor (e.g., a dosage sufficient to lower the serum calcium level to below 10 mg/dl). Typical dosage would be 1 mg to 10 mg of the 35 antibody or peptide per kg body weight per day.

Treatment may be repeated as necessary for long term maintenance of acceptable calcium levels (i.e., levels < 10.1 mg/dl). This may be necessary for acute treatment of an underlying disease condition triggering bypercalcemia; or it may used, e.g., for chronic

treatment of conditions such as osteoporosis.

In another application, the compounds of the invention which have been characterized, according to the methods of the invention, to be agonists are used

10 therapeutically to treat hypocalcemia: e.g., that resulting from the partial or complete surgical removal of the parathyroid glands. Agonists may be formulated in a suitable carrier (e.g., physiological saline) and are preferably administered intravenously in a dosage that

15 causes a rise in serum calcium to an acceptable level (i.e., approximately 8 mg/dl). A useful dosage range would be 1 ng to 10 mg of the agonist per kg body weight

Treatment may be repeated as necessary to

maintain suitable serum calcium levels; long term 20 treatment may be necessary for patients who have undergone parathyroid gland removal.

The nucleic acids of the invention may also be used therapeutically. Oligonucleotides which are antisense to PTH receptor mRNA (or nucleic acid constructs which express RNA that is antisense to PTH receptor mRNA) may be utilized as an anticancer therapy. This approach is useful, e.g., for hypercalcemias resulting from a genomic rearrangement or amplification which increases the amount or activity of PTH receptor, PTH or PTHrP. The method would involve introduction of the antisense oligonucleotide into the tumor cells in vivo. The antisense strand hybridizes with endogenous PTH receptor mRNA, interfering with translation of the protein, thereby reducing production of PTH receptor in such cells, and reducing PTH/PTHrP-associated neoplastic

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growth. Methods for antisense design and introduction into host cells are described, for example, in Weinberg et al., U.S. Patent No. 4,740,463, herein incorporated by The biochemical characterization of the OKreference. 5 H, OK-O and R15B PTH/PTHrP receptors of the invention demonstrate that the two transduction pathways now known to be triggered by the interaction of PTH with its receptor are distinct and may be separated. predicted amino acid sequences of these receptors 10 indicate that OK-H, which does not appear to activate inositol phosphate metabolism to any detectable degree, is 70 amino acids shorter at the carboxy-terminus than OK-O or R15B. By using the sequences of the invention and the information disclosed herein, one could clone and 15 then alter (e.g. by site-directed mutagenesis) PTH/PTHrP receptor genes from any species to generate PTH/PTHrP receptors which do not activate phospholipase C. This could potentially allow the separation of different PTHmediated actions, including bone resorption and bone 20 formation, and could of great importance for the treatment of various bone disorders such as osteoporosis.

Nucleic acids of the invention which encode a PTH receptor may also be linked to a selected tissue-specific promoter and/or enhancer and the resultant hybrid gene introduced, by standard methods (e.g., as described by Leder et al., U.S. Patent No. 4,736,866, herein incorporated by reference), into an animal embryo at an early developmental stage (e.g., the fertilized oocyte stage), to produce a transgenic animal which expresses elevated levels of PTH receptor in selected tissues (e.g., the osteo calcin promoter for bone). Such promoters are used to direct tissue-specific expression of the PTH receptor in the transgenic animal. The form of PTH receptor utilized can be one which encodes a PTH receptor similar to that of the animal species used, or

it can encode the PTH receptor homolog of a different species. In one particular example, transgenic chickens are engineered to express the PTH receptor from a promoter which directs high-level expression in chicken oviducts. Such an animal is expected to produce eggs with higher calcium content, and thus harder shells.

Other Embodiments

Other embodiments are within the following claims. For example, the nucleic acid of the invention includes genes or cDNAs or RNAs originally isolated from any vertebrate species, including birds or mammals such as marsupials, rodents, or humans. The high degree of homology demonstrated for the PTH receptors from such diverse species as opossum, rat, and human indicates that the methods of isolating PTH receptors disclosed herein will be broadly applicable to the isolation of related cell receptors from a wide variety of species.

PCT/US92/02821

COMPUTER SUBMISSION OF DNA AND AMINO ACID SEQUENCES

(1) GENERAL INFORMATION:

(i) APPLICANT:

Segre, Gino V.

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Juppner, Harald Potts, John T., Jr. Schipani, Ernestina

(ii) TITLE OF INVENTION:

PARATHYROID HORMONE RECEPTOR AND DNA

ENCODING SAME

(iii) NUMBER OF SEQUENCES:

(iv) CORRESPONDENCE ADDRESS:

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(D) STATE:

Massachusetts

(E) COUNTRY:

U.S.A.

(F) ZIP:

02110-2804

(v) COMPUTER READABLE FORM:

(A) MEDIUM TYPE:

3.5" Diskette, 1.44 Mb storage

(B) COMPUTER:

IBM PS/2 Model 50Z or 55SX

(C) OPERATING SYSTEM:

IBM P.C. DOS (Version 3.30)

(D) SOFTWARE:

WordPerfect (Version 5.0)

- (vi) CURRENT APPLICATION DATA:
- (A) APPLICATION NUMBER:
- (B) FILING DATE:
- (C) CLASSIFICATION:
- (vii) PRIOR APPLICATION DATA:

(A) APPLICATION NUMBER:

07/681,702

(B) FILING DATE:

April 5, 1991

(viii) ATTORNEY/AGENT INFORMATION:

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- 51 -

(C) TELEX:

200154

(2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 1:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 1862
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: double
- (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQUENCE ID NO: 1:

(xi) SEQUE	NCE DESCRIPTI	ON: SEQUENC	E ID NO: 1:	
TGGGCACAGC CACCC GTGGCCCCGT TGGAC			ATG GGA GCG	60
TCG CAC AGC CTT Ser His Ser Leu 10				163
TAC GCA CTG GTG Tyr Ala Leu Val	Asp Ala Asp A			211
ATT CTT CTG CGC . Ile Leu Leu Arg . 40				259
GTC CTC AGG GTC Val Leu Arg Val 55				307
AGG TCT GCA AAG . Arg Ser Ala Lys		lu Lys Pro		355
CAG GCA GAG GAG GIn Ala Glu Glu 90	Ser Arg Glu V			403
GGC TTC TGC CTA Gly Phe Cys Leu 105	Pro Glu Trp A			451
GTG CCC GGC AAG Val Pro Gly Lys 120				 499
TTC AAC CAC AAA Phe Asn His Lys 135				547

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											GCG					595
Trp	Glu	Leu	Val	Pro	Gly	Asn	Asn	Arg	Thr	Trp	Ala	Asn	Tyr	Ser	Glu	
				15	55				16	50				16	55	
																643
TGT	GTC	AAG	TTT	CTG	ACC	AAC	GAG	ACC	CGG	GAA	CGG	GAA	GTC	TTT	GAT	043
Сув	Val	Lys			Thr	Asn	GIU			GIU	Arg	GIU	18		мвр	
			17	70				17	75				10			
ccc	CTC	CCA	ATC	A TIC	TAC	እ ርጥ	стс	GGC	TAC	TCC	ATC	TCT	CTG	GGC	TCC	691
Ara	T.em	Glv	Met	Tle	Tvr	Thr	Val	Glv	Tvr	Ser	Ile	Ser	Leu	Gly	Ser	
my	Dec	185			-,-		190		- 4			195		-		
											AGG					739
Leu	Thr	Val	Ala	Val	Leu	Ile	Leu	Gly	Tyr	Phe	Arg	Arg	Leu	His	Сув	
	200					205					210					
																787
ACC	CGA	AAC	TAC	ATT	CAC	ATG	CAT	CTC	TTC	GTG	TCC	TTT	ATG	CTC	CGG	787
	Arg	Asn	Tyr	Ile		Met	His	Leu	Pne		Ser	Pne	Met	Leu		
215					220					225					230	
COTT	CTA	ACC	እጥሮ	ሙውር	እ ጥር	ממג	CAT	CCT	стс	СТС	TAC	TCG	GGG	GTT	TCC	835
											Tyr					
VIG	AGT	361	110		35	_, _				10	-1-		2	24		
				-										•		
ACA	GAT	GAA	ATC	GAG	CGC	ATC	ACC	GAG	GAG	GAG	CTG	AGG	GCC	TTC	ACA	883
											Leu					
	_		250					255					260			
																021
GAG	CCT	CCC	CCT	GCT	GAC	AAG	GCG	GGT	TTT	GTG	GGC	TGC	AGA	GTG	GCG	931
Glu	Pro		Pro	Ala	Asp	Lys	_	Gly	Phe	Val	Gly		Arg	vaı	AIG	
		265					270					275				
CTD	3.00	CTC	ጥጥር		ጥልሮ	ጥጥር	CTG	ACC	ACC	AAC	TAC	TAC	TGG	ATC	CTG	979
											Tyr					
141	280	741			-1-	285					290		•			
GTG	GAA	GGC	CTC	TAC	CTT	CAC	AGC	CTC	ATC	TTC	ATG	GCT	TTT	TTC	TCT	1027
Val	Glu	Gly	Leu	Tyr	Leu	His	Ser	Leu	Ile	Phe	Met	Ala	Phe	Phe	Ser	
295		_		-	300					305					310	
GAG	AAA	AAG	TAT	CTC	TGG	GGT	TTC	ACA	TTA	TTT	GGC	TGG	GGC	CTC	CCT	1075
Glu	Lys	Lys	Tyr			Gly	Phe	Thr			Gly	Trp	GTĀ			
				3	15				3.	20				34	25	
ccc	CMC	ተነጥጥ	ርሞሶ	COP	GTG	ጥርር	GTG	ACC	ርጥር	AGG	GCT	ACA	CTG	GCC	AAC	1123
212	A=1	Pho	Val	Ala	Val	Trn	Val	Thr	Val	Aro	Ala	Thr	Leu	Ala	Asn	
viq	447	1116	330	mra	*44	2		335		3			340			
			-55													
ACT	GAG	TGC	TGG	GAC	CTG	AGT	TCG	GGG	AAT	AAG	AAA	TGG	ATC	ATA	CAG	1171
Thr	Glu	Сув	Trp	Asp	Leu	Ser	Ser	Gly	Asn	Lys	Lys	Trp	Ile	Ile	Gln	
		345	-	-			350					355				

										TTT Phe						1219
vaı	360	116	Leu	WIG	MIG	365	Val	Val	ABII	Pne	370	Leu	Pne	IIe	ABN	
										GAG						1267
375	Ile	Arg	Val	Leu	Ala 380	Thr	Lys	Leu	Arg	Glu 385	Thr	Asn	Ala	Gly	_	
															390	
										CTG						1315
Сув	Авр	The	Arg		GIN 95	Tyr	Arg	гÀа	Leu 40	Leu	ràs	ser	Thr	Leu 40		
										_					-	
										GTC						1363
Leu	Met	Pro			Gly	Val	His	_		Val	Phe	Met			Pro	
			4.	lO				41	15				42	20		
TAC	ACA	GAA	GTA	TCA	GGG	ATT	CTT	TGG	CAA	GTC	CAA	ATG	CAC	TAT	GAA	1411
Tyr	Thr		Val	Ser	Gly	Ile		Trp	Gln	Val	Gln	Met	His	Tyr	Glu	
		425					430					435				
ATG	CTC	TTC	AAT	TCA	TTC	CAG	GGA	TTT	TTC	GTT	GCC	ATT	ATA	TAC	TGT	1459
Met		Phe	Asn	Ser	Phe		Gly	Phe	Phe	Val		Ile	Ile	Tyr	Сув	
	440					445					450					
TTC	TGC	AAT	GGA	GAG	GTA	CAA	GCA	GAG	ATC	AAG	AAG	TCA	TGG	AĞC	CGA	1507
	Сув	Asn	Gly	Glu		Gln	Ala	Glu	Ile	Lys	Lys	Ser	Trp	Ser	_	
455					460					465					470	
TGG	ACC	CTG	GCC	TTG	GAC	TTC	AAG	CGG	AAG	GCC	CGG	AGT	GGC	AGC	AGT	1555
Trp	Thr	Leu	Ala		-	Phe	Lys	Arg		Ala	Arg	Ser	Gly			
				47	75				48	30				.48	35	
ACC	TAC	AGC	TAT	GGC	CCC	ATG	GTG	TCA	CAT	ACA	AGT	GTC	ACC	AAT	GTG	1603
Thr	Tyr	Ser		Gly	Pro	Met	Val		His	Thr	Ser	Val		Asn	Val.	
			490					495					500			
GGA	CCT	CGA	GGG	GGC	TGG	CCT	TGT	ccc	TCA	GCC	CTC	GAC	TAGO	CTCCI	rgg	1652
Gly	Pro	-	Gly	Gly	Trp	Pro	-	Pro	Ser	Ala	Leu	-	•			
		505					510					515				
															TTCCAT	
															STATCT	
					rg ac				-GGA	ACAG	CCCC	CTC	JAC 7	CCTC	GAGGA	1832 1862
JUNE	'UAU	t var			2 7 G	2006	* 1 14 1 (•								1007

(2)	INFO	RMAT	MOI	FOR	SEQU	ENCE	IDE	NTIF	PICAT	CION	NUMB	ER:	2	:		
	(i	.) SE	QUEN	ICE C	CHARA	CTEF	ISTI	cs:								
		(• •	YPE:					1863 nucl sing	eic ;le	acid	I				
	(3	:i) S	EQUE	NCE	DESC	RIPT	NOI:	SEC	QUENC	E II	NO:	2:				
TGGG	CACA	AGC C	CACCO	TGTI	rg gi	AGTO	CAGG	GGC	CAGC	CCA	CTGA	GCTG	GC A	TATC	AGCTG	60
GTGG	ccc	GT 1	OADDT	CTCGG	SC CC	TAGG	GAAC	GGC	CGCC	ATC Met	: Gly	GCG Ala	CCC Pro	CGG Arg	ATC	115
TCG Ser	CAC His	AGC Ser	CTT Leu 10	Ala	TTG Leu	CTC Leu	CTC Leu	TGC Cys	Сув	TCC Ser	GTG Val	CTC Leu	AGC Ser 20	TCC Ser	GTC Val	163
TAC Tyr	GCA Ala	CTG Leu 25	GTG Val	GAT Asp	GCC Ala	GAT Asp	GAT Asp 30	GTC Val	ATA Ile	ACG Thr	AAG Lys	GAG Glu 35	GAG Glu	CAG Gln	ATC Ile	211
ATT Ile	CTT Leu 40	CTG Leu	CGC Arg	AAT Asn	GCC Ala	CAG Gln 45	GCC Ala	CAG Gln	TGT Cys	GAG Glu	CAG Gln 50	CGC Arg	CTG Leu	AAA Lys	GAG Glu	259
GTC Val 55	CTC Leu	AGG Arg	GTC Val	CCT Pro	GAA Glu 60	CTT Leu	GCT Ala	GAA Glu	TCT Ser	GCC Ala 65	TA8	GAC Asp	TGG Trp	ATG Met	TCA Ser 70	307
AGG Arg	TCT Ser	GCA Ala	AAG Lys	ACA Thr	Lys	AAG Lys	GAG Glu	AAA Lys	CCT Pro	Ala	GAA Glu	AAG Lys	CTT Leu	TAT Tyr 85	Pro	355
CAG Gln	GCA Ala	GAG Glu	GAG Glu 9	Ser	AGG Arg	GAA Glu	GTT Val	TCT Ser	Asp	AGG Arg	AGC Ser	CGG Arg	Leu	CAG Gln 00	GAT Asp	403
GGC Gly	TTC Phe	TGC Cys 105	Leu	CCT Pro	GAG Glu	TGG Trp	GAC Asp 110	AAC Asn	ATT Ile	GTG Val	TGC Cys	TGG Trp 115	CCT Pro	GCT Ala	GGA Gly	451
GTG Val	CCC Pro	GGC Gly	AAG Lys	GTG Val	GTG Val	GCC Ala	GTG Val	CCC Pro	TGC Cys	CCC Pro	GAC Asp	TAC Tyr	TTC Phe	TAC Tyr	GAC Asp	499

125

120

										TGT Cys 145						547
				Pro					Thr	TGG Trp 50					Glu	595
				. – –						GAA Glu						643
										TCC Ser						691
										TTT Phe						739
										GTG Val 225						787
										CTC Leu						835
				2.3	35	_			24	10	_			24	15	
		Glu		GAG	CGC				GAG	GAG Glu				TTC	ACA	883
Thr	Asp	Glu CCC	Ile 250 CCT	GAG Glu GCT	CGC Arg	Ile AAG	Thr	Glu 255 GGT	GAG Glu TTT	GAG	Leu GGC	Arg TGC	Ala 260 AGA	TTC Phe GTG	ACA Thr	931
Thr GAG Glu GTA	Asp CCT Pro	Glu CCC Pro 265	Ile 250 CCT Pro	GAG Glu GCT Ala	CGC Arg GAC Asp	Ile AAG Lys TTC	Thr GCG Ala 270 CTG	Glu 255 GGT Gly ACC	GAG Glu TTT Phe	GAG Glu GTG	CGC Gly	TGC Cys 275	Ala 260 AGA Arg	TTC Phe GTG Val	ACA Thr GCG Ala	
GAG Glu GTA Val	CCT Pro ACC Thr 280	Glu CCC Pro 265 GTC Val	CCT Pro	GAG Glu GCT Ala CTT Leu	CGC Arg GAC Asp TAC Tyr	AAG Lys TTC Phe 285	Thr GCG Ala 270 CTG Leu AGC	Glu 255 GGT Gly ACC Thr	GAG Glu TTT Phe ACC Thr	GAG Glu GTG Val	GGC Gly TAC Tyr 290	TGC Cys 275 TAC Tyr	Ala 260 AGA Arg TGG Trp	TTC Phe GTG Val ATC Ile	ACA Thr GCG Ala CTG Leu	931
GAG Glu GTA Val GTG Val 295	ABP CCT Pro ACC Thr 280 GAA Glu	CCC Pro 265 GTC Val GGC Gly	Ile 250 CCT Pro TTC Phe CTC Leu	GAG Glu GCT Ala CTT Leu TAC Tyr	CGC Arg GAC Asp TAC Tyr CTT Leu 300	AAG Lys TTC Phe 285 CAC His	Thr GCG Ala 270 CTG Leu AGC Ser	Glu 255 GGT Gly ACC Thr CTC Leu	GAG Glu TTT Phe ACC Thr ATC Ile	GAG Glu GTG Val AAC Asn	GGC Gly TAC Tyr 290 ATG Met	TGC Cys 275 TAC Tyr GCT Ala	Ala 260 AGA Arg TGG Trp TTT Phe	TTC Phe GTG Val ATC Ile TTC Phe	ACA Thr GCG Ala CTG Leu TCT Ser 310	931 979

ACT Thr	GAG Glu	TGC Cys 345	TGG Trp	GAC Asp	CTG Leu	AGT Ser	TCG Ser 350	GGG Gly	AAT Asn	AAG Lys	AAA Lys	TGG Trp 355	ATC Ile	ATA Ile	CAG Gln	1171
GTG Val	CCC Pro 360	ATC Ile	CTG Leu	GCA Ala	GCT Ala	ATT Ile 365	GTG Val	GTG Val	AAC Asn	TTT Phe	ATT Ile 370	CTT Leu	TTT Phe	ATC Ile	AAT Asn	1219
ATA Ile 375	ATC Ile	AGA Arg	GTC Val	CTG Leu	GCT Ala 380	ACT Thr	AAA Lys	CTC Leu	CGG Arg	GAG Glu 385	ACC Thr	AAT Asn	GCA Ala	GGG Gly	AGA Arg 390	1267
TGT Cys	GAC Asp	ACG Thr	AGG Arg	Gln	CAG Gln 95	TAT Tyr	AGA Arg	AAG Lys	CTG Leu 40	Leu	AAG Lys	TCC Ser	ACG Thr	Leu	GTC Val 05	1315
CTC Leu	ATG Met	CCG Pro	CTA Leu 410	TTT Phe	GGG Gly	GTG Val	CAC His	TAC Tyr 415	ATC Ile	GTC Val	TTC Phe	ATG Met	GCC Ala 420	ACG Thr	CCG Pro	1363
TAC Tyr	ACA Thr	GAA Glu 425	GTA Val	TCA Ser	GGG Gly	ATT Ile	CTT Leu 430	TGG Trp	CAA Gln	GTC Val	CAA Gln	ATG Met 435	CAC His	TAT Tyr	GAA Glu	1411
ATG Met	CTC Leu 440	TTC Phe	AAT	TCA Ser	TTC Phe	CAG Gln 445	GGA Gly	TTT Phe	TTC Phe	GTT Val	GCC Ala 450	ATT Ile	ATA Ile	TAC Tyr	TGT Cys	1459
TTC Phe 455	TGC Cys	AAT Asn	GGA Gly	GAG Glu	GTA Val 460	CAA Gln	GCA Ala	GAG Glu	ATC Ile	AAG Lys 465	AAG Lys	TCA Ser	TGG Trp	AGC Ser	CGA Arg 470	1507
TGG Trp	ACC Thr	CTG Leu	GCC Ala	Leu	GAC Asp 75	TTC Phe	AAG Lys	CGG Arg	Lys	GCC Ala BO	CGG Arg	AGT Ser	GGC Gly	Ser	AGT Ser 85	1555
ACC Thr	TAC Tyr	AGC Ser	TAT Tyr 490	Gly	CCC	ATG Met	GTG Val	TCA Ser 495	CAT His	ACA Thr	AGT Ser	GTC Val	ACC Thr 500	AAT Asn	GTG Val	1603
GGA Gly	CCT Pro	CGA Arg 505	Gly	GGG	CTG Leu	GCC Ala	TTG Leu 510	Ser	CTC	AGC Ser	CCT Pro	CGA Arg 515	CTA Leu	GCT Ala	CCT Pro	1651
GGG Gly	GCT Ala 520	GGA Gly	GCC Ala	AGT Ser	GCC	AAT Asn 525	Gly	CAT	CAC His	CAG Gln	TTG Leu 530	Pro	GGC	TAT	GTG Val	1699
AAG Lys 535	CAT His	GGT Gly	TCC Ser	ATT	TCT Ser 540	Glu	AAC	TCA Ser	TTG Leu	CCT Pro 545	Ser	TCT	GGC	CCA Pro	GAG Glu 550	1747

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CCT																
															GAG	1795
Pro	Gly	Thr	Lys	Asp	Asp	Gly	Tyr	Leu	Asn	Gly	Ser	Gly	Leu	Tyr	Glu	
				5	5 5				5	60				5	65	
003	3.00												•			
Dro	MOT	GTT	GGG	GAA	CAG	CCC	CCT	CCA	CTC	CTG	GAG	GAG	GAG	AGA	GAG	1843
PIO	Mec	Val	570	GIU	GIN	PFO			Leu	Leu	GIU	Glu		Arg	Glu	
			570				•	575					580			
ACA	GTC	ATG	TCAC	יכראי	י ידעי	,		٠								1000
	Val		2 0	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	ını (•									•	1863
		585														
(2)	INF	ORMA:	CION	FOR	SEQ	JENCI	IDI	ENTI	FICA'	TION	NUM	BER:		3:		
	(:	i) SI	QUE	ICE (CHAR	CTE	RIST	ICS:								
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	/-	xi) S	EOH	יאריבי	חדיכו	ים ד מי	TON	CE/	NETTO NE	7D T1	. NO.					
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GGC	GGGG	cc c	CGGC	GGC	A GO	TCG	AGGC	CGC	cccc	CGGC	TGCC	ccc	AGG (ZAČGO	CGGCCC	60
TAG	GCGG:	rgg (G AI	'G GC	G GC	C G	C CC	G A	rc G	CA CO	C AC	C C	rg g	CG C	rc rc	108
										la Pi						100
				1				•	5					10		•
														٠		
		TGC														156
Leu	Leu	Cys	Сув	Pro	Val	Leu		Ser	Ala	Tyr	Ala	Leu	Val	Acr	Ala	
		15					20			-				veħ		
CNC	CAM									_		25		veb		,
GAC	GWI	CMC	mmm	200	222	~~~		222	.			25		_		
Aan	Aen	GTC Val	TTT	ACC	AAA	GAG	GAA	CAG	ATT	TTC	CTG	25 CTG	CAC	CGT	GCC	204
Asp	Asp	GTC Val	TTT Phe	ACC Thr	AAA Lys	Glu	GAA	CAG Gln	ATT Ile	TTC	CTG Leu	25 CTG	CAC	CGT	GCC	
Asp	Asp 30	GTC Val	TTT Phe	ACC Thr	AAA Lys	GAG Glu 35	GAA	CAG Gln	ATT Ile	TTC	CTG	25 CTG	CAC	CGT	GCC	
	Asp 30	Val	Phe	Thr	Lys	Glu 35	GAA Glu	Gln	Ile	TTC Phe	CTG Leu 40	25 CTG Leu	CAC His	CGT Arg	GÇC Ala	204
CAG	Asp 30 GCG	Val CAA	Phe TGT	Thr	Lys AAG	Glu 35 CTG	GAA Glu CTC	Gln AAG	Ile GAA	TTC Phe GTT	CTG Leu 40 CTG	25 CTG Leu CAC	CAC His	CGT Arg	GCC Ala	
CAG	Asp 30 GCG	Val	Phe TGT	Thr	Lys AAG	Glu 35 CTG	GAA Glu CTC	Gln AAG	Ile GAA	TTC Phe GTT	CTG Leu 40 CTG	25 CTG Leu CAC	CAC His	CGT Arg	GCC Ala	204
CAG Gln	Asp 30 GCG	Val CAA	Phe TGT	Thr	Lys AAG Lys	Glu 35 CTG	GAA Glu CTC	Gln AAG	Ile GAA	TTC Phe GTT Val	CTG Leu 40 CTG	25 CTG Leu CAC	CAC His	CGT Arg	GCC Ala GCC Ala	204
CAG Gln 45	Asp 30 GCG Ala	Val CAA	TGT Cys	Thr GAC Asp	AAG Lys 50	Glu 35 CTG Leu	GAA Glu CTC Leu	Gln AAG Lys	Ile GAA Glu	TTC Phe GTT Val 55	CTG Leu 40 CTG Leu	25 CTG Leu CAC His	CAC His ACA Thr	CGT Arg GCA Ala	GCC Ala GCC Ala 60	204
CAG Gln 45 AAC	Asp 30 GCG Ala ATA	Val CAA Gln	TGT Cys	Thr GAC Asp TCA	AAG Lys 50	Glu 35 CTG Leu AAG	GAA Glu CTC Leu	Gln AAG Lys TGG	Ile GAA Glu ACA	TTC Phe GTT Val 55	CTG Leu 40 CTG Leu	25 CTG Leu CAC His	CAC His ACA Thr	CGT Arg GCA Ala	GCC Ala GCC Ala 60	204
CAG Gln 45 AAC	Asp 30 GCG Ala ATA	CAA Gln ATG	TGT Cys	Thr GAC Asp TCA Ser	AAG Lys 50	Glu 35 CTG Leu AAG	GAA Glu CTC Leu	Gln AAG Lys TGG	GAA Glu ACA Thr	TTC Phe GTT Val 55	CTG Leu 40 CTG Leu	25 CTG Leu CAC His	CAC His ACA Thr	CGT Arg GCA Ala TCA Ser	GCC Ala GCC Ala 60	204
CAG Gln 45 AAC ABN	Asp 30 GCG Ala ATA Ile	CAA Gln ATG Met	TGT Cys GAG Glu	GAC Asp TCA Ser	AAG Lys 50 GAC Asp	Glu 35 CTG Leu AAG Lys	GAA Glu CTC Leu GGC	Gln AAG Lys TGG Trp	GAA Glu ACA Thr	TTC Phe GTT Val 55 CCA Pro	CTG Leu 40 CTG Leu GCA Ala	25 CTG Leu CAC His	CAC His ACA Thr ACG Thr	CGT Arg GCA Ala TCA Ser	GCC Ala GCC Ala 60 GGG Gly	204
CAG Gln 45 AAC ABN	Asp 30 GCG Ala ATA Ile	CAA Gln ATG Met	TGT Cys GAG Glu	GAC Asp TCA Ser	AAG Lys 50 GAC Asp	Glu 35 CTG Leu AAG Lys	GAA Glu CTC Leu GGC Gly	AAG Lys TGG Trp	GAA Glu ACA Thr	TTC Phe GTT Val 55 CCA Pro 70	CTG Leu 40 CTG Leu GCA Ala	25 CTG Leu CAC His TCT Ser	CAC His ACA Thr ACG Thr	CGT Arg GCA Ala TCA Ser	GCC Ala GCC Ala 60 GGG Gly 75	204
CAG Gln 45 AAC ABN	Asp 30 GCG Ala ATA Ile	CAA Gln ATG Met	TGT Cys GAG Glu AAA Lys	GAC Asp TCA Ser GAG Glu	AAG Lys 50 GAC Asp	Glu 35 CTG Leu AAG Lys	GAA Glu CTC Leu GGC Gly	AAG Lys TGG Trp GGA Gly	GAA Glu ACA Thr AAG Lys	TTC Phe GTT Val 55 CCA Pro 70	CTG Leu 40 CTG Leu GCA Ala	25 CTG Leu CAC His TCT Ser	CAC His ACA Thr ACG Thr	CGT Arg GCA Ala TCA Ser	GCC Ala GCC Ala 60 GGG Gly 75	204 252 300
CAG Gln 45 AAC ABN	Asp 30 GCG Ala ATA Ile	CAA Gln ATG Met	TGT Cys GAG Glu	GAC Asp TCA Ser GAG Glu	AAG Lys 50 GAC Asp	Glu 35 CTG Leu AAG Lys	GAA Glu CTC Leu GGC Gly	AAG Lys TGG Trp	GAA Glu ACA Thr AAG Lys	TTC Phe GTT Val 55 CCA Pro 70	CTG Leu 40 CTG Leu GCA Ala	25 CTG Leu CAC His TCT Ser	CAC His ACA Thr ACG Thr	CGT Arg GCA Ala TCA Ser	GCC Ala GCC Ala 60 GGG Gly 75	204 252 300
CAG Gln 45 AAC ABN AAG Lys	Asp 30 GCG Ala ATA Ile CCC Pro	CAA Gln ATG Met AGG Arg	TGT Cys GAG Glu AAA Lys	GAC Asp TCA Ser GAG Glu	AAG Lys 50 GAC Asp 55	Glu 35 CTG Leu AAG Lys GCA Ala	GAA Glu CTC Leu GGC Gly TCG Ser	AAG Lys TGG Trp GGA Gly	GAA Glu ACA Thr AAG Lys	TTC Phe GTT Val 55 CCA Pro 70 TTC Phe	CTG Leu 40 CTG Leu GCA Ala TAC	25 CTG Leu CAC His TCT Ser CCT Pro	CAC His ACA Thr ACG Thr GAG Glu	CGT Arg GCA Ala TCA Ser TCT	GCC Ala GCC Ala 60 GGG Gly 75 AAA Lys	204 252 300
CAG Gln 45 AAC ABN AAG Lys	ASP 30 GCG Ala ATA Ile CCC Pro	CAA Gln ATG Met	TGT Cys GAG Glu AAA Lys 80 GAC	GAC Asp TCA Ser GAG Glu	AAG Lys 50 GAC Asp 55 AAG Lys	Glu 35 CTG Leu AAG Lys GCA Ala	GAA Glu CTC Leu GGC Gly TCG Ser	AAG Lys TGG Trp GGA Gly 85	GAA Glu ACA Thr AAG Lys	TTC Phe GTT Val 55 CCA Pro 70 TTC Phe	CTG Leu 40 CTG Leu GCA Ala TAC Tyr	25 CTG Leu CAC His TCT Ser CCT Pro	CAC His ACA Thr ACG Thr GAG Glu 90	CGT Arg GCA Ala TCA Ser TCT Ser	GCC Ala GCC Ala 60 GGG Gly 75 AAA Lys	204 252 300

	TGG Trp								444
	GCA Ala								492
	GCC Ala	Tyr			Arg			Glu	540
	CAC His 160								588
	AAT Asn								636
	ACC Thr								684
	ATC Ile								732
	ATG Met	His			Phe			Ala	780
	AAG Lys 240								828
	CTC Leu								876
	GCC Ala								924
	TTC Phe								972
	CTG Leu	Tyr			Ile			Phe	1020

				TTT GGC TG Phe Gly Tr		
Ala Val Pl			l Gly Val	AGA GCA AC Arg Ala Th 34	r Leu Ala	
				AAG AAG TG Lys Lys Tr 360		
				TTC ATC CT Phe Ile Let 375		
				GAG ACC AA		
				CTC AGG TC		
Leu Val Pr			Tyr Thr	GTC TTC ATC Val Phe Me 42	t Ala Leu	
				ATC CAG ATG		
	he Asn Ser		y Phe Phe	GTT GCC AT Val Ala Il 55	e Ile Tyr	
				AGG AAG TC Arg Lys Se		
Trp Thr Le			a Arg Lys	GCA CGA AG Ala Arg Se 49	r Gly Ser	
				ACG AGT GT Thr Ser Va 505		
GGC CCC CC	GT GCA GGA	CTC AGC CT	C CCC CTC	AGC CCC CG	C CTG CCT	CCT 1644

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				Gly					Pro					CCA Pro		1692
			Thr				Ţhr		Pro				Ala	GTT Val 55		1740
														GAG Glu		1788
														TGG Trp		1836
	GTC Val 590		TGA	CTGG	GCA (CTAGO	GGGG	CT AC	BACTO	GCTG	G CC	rggg(CACA			1885
AAG	ATAA	CAA I	AAGG	AAAA!	rg G	AAGT	TTTC GACC	AA C	CAG	AGAA	GAA	GAA(GAT (CTGG! GTTT:	ACCAGG FGCAGG	1945 2005 2051

What is claimed is:

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Claims

- 1. Isolated DNA comprising a DNA sequence
- 2 encoding a cell receptor of a vertebrate animal, said
- 3 receptor having an amino acid sequence with at least 30%
- 4 identity to the amino acid sequence shown in FIG. 3.
- 1 2. The isolated DNA of claim 1, wherein said
- 2 DNA sequence encodes substantially all of the amino acid
- 3 sequence shown in FIG. 1 (SEQ. ID NO. 1).
- 1 3. The isolated DNA of claim 1, wherein said
- 2 DNA sequence encodes substantially all of the amino acid
- 3 sequence shown in FIG. 3 (SEQ. ID NO. 3).
- 1 4. The isolated DNA of claim 1, said isolated
- 2 DNA being (8A6), deposited with the ATCC and designated
- 3 ATCC Accession No. 68570.
- 5. The isolated DNA of claim 1, wherein said
- 2 DNA sequence encodes substantially all of the amino acid
- 3 sequence shown in Fig. 6 (SEQ. ID. NO. 4).
- 1 6. The isolated DNA of claim 1, wherein said
- 2 DNA sequence hybridizes to the DNA sequence shown in Fig.
- 3 1 (SEQ. ID NO. 1).
- The isolated DNA of claim 1, wherein said
- 2 DNA sequence hybridizes to the DNA sequence shown in Fig.
- 3 3 (SEQ. ID NO. 3).
- 1 8. The isolated DNA of claim 1, wherein said
- 2 DNA sequence hybridizes to the DNA sequence shown in Fig.
- 3 6 (SEQ. ID NO. 4).

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- 9. A purified preparation of a vector, said
- 2 vector comprising a DNA sequence encoding a parathyroid
- 3 hormone receptor.
- 1 10. A cell containing the isolated DNA of claim
- 2 1.
- 1 11. The cell of claim 10, wherein said cell is
- 2 capable of expressing said cell receptor from said
- 3 isolated DNA.
- 1 . An essentially homogenous population of
- 2 cells, each of which comprises the isolated DNA of claim
- 3 1.

- 1 13. Isolated DNA comprising a DNA sequence
- 2 encoding a polypeptide capable of binding parathyroid
- 3 hormone or parathyroid-hormone-related protein.
- 1 14. A method for producing a polypeptide, said
- 2 method comprising:
- 3 providing a cell comprising isolated DNA
- 4 encoding a parathyroid hormone receptor or a fragment
- 5 thereof; and
- 6 culturing said cell under conditions
- 7 permitting expression of a polypeptide from said DNA.
- 1 15. A single-stranded DNA comprising a portion
- 2 of a parathyroid hormone receptor gene, said portion
- 3 being at least 18 nucleotides long.
- 1 16. The single-stranded DNA of claim 15, wherein
- 2 said portion is less than all of said parathyroid hormone
- 3 receptor gene.

- 1 17. The single-stranded DNA of claim 15, wherein
- 2 said DNA is detectably labeled.
- 1 18. A single-stranded DNA comprising a portion
- 2 of a parathyroid hormone receptor cDNA, said portion
- 3 being at least 18 nucleotides long.
- 1 19. The single-stranded DNA of claim 18, wherein
- 2 said DNA is antisense.
- 1 20. Parathyroid hormone receptor produced by
- 2 expression of a recombinant DNA molecule encoding a
- 3 parathyroid hormone receptor.
- 1 21. An essentially purified preparation of the
- 2 parathyroid hormone receptor of claim 20.
- 1 . 22. An essentially purified preparation of the
- 2 parathyroid receptor produced by the expression of the
- 3 DNA of claim 5.
- 23. A polypeptide comprising at least six amino
- 2 acids and less than the complete amino acid sequence of a
- 3 parathyroid hormone receptor, said polypeptide capable of
- 4 binding parathyroid hormone or parathyroid hormone-
- 5 related protein.
- 1 24. The polypeptide of claim 23, wherein said
- 2 parathyroid hormone receptor is a human parathyroid
- 3 receptor.
- 1 25. The polypeptide of claim 23, wherein said
- 2 fragment comprises
- 3 (a) TNETREREVFDRLGMIYTVG,
- 4 (b) YLYSGFTLDEAERLTEEEL,

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5	(c) [.]	VTFFLYFLATNYYWILVEG,
. 6	(d)	Y-RATLANTGCWDLSSGHKKWIIQVP,
7	(e)	PYTEYSGTLWQIQMHYEM,
8	(f)	DDVFTKEEQIFLLHRAQA,
9	(g)	FFRLHCTRNY,
10	(h)	EKKYLWGFTL,
11	(i)	VLATKLRETNAGRCDTRQQYRKLLK, or
12	(j)	a fragment of (a) - (i) which is capable of
13	binding par	athyroid hormone or parathyroid hormone-

- 1 26. A therapeutic composition comprising, in a
- 2 pharmaceutically-acceptable carrier, (a) a parathyroid
- 3 hormone receptor or (b) a polypeptide comprising a
- 4 fragment of said receptor.

related protein.

- 27. An antibody capable of forming an immune complex with a parathyroid hormone receptor.
- 1 28. A therapeutic composition comprising the
- 2 antibody of claim 27 and a pharmaceutically-acceptable
- 3 carrier.

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- 1 29. A method of reducing the level of calcium in
- 2 the blood of a mammal, which method comprises
- 3 administering the therapeutic composition of claim 26 to
- 4 said mammal in a dosage effective to inhibit activation
- 5 by parathyroid hormone or parathyroid hormone-related
- 6 protein of a parathyroid hormone receptor of said mammal.
- 1 30. A method of reducing the level of calcium in
- 2 the blood of a mammal, which method comprises
- 3 administering the therapeutic composition of claim 28 to
- 4 said mammal in a dosage effective to inhibit activation

- 5 by parathyroid hormone or parathyroid hormone-related
- 6 protein of a parathyroid hormone receptor of said mammal.
- 1 31. A method for identifying a compound capable
- 2 of competing with a parathyroid hormone for binding to a
- 3 parathyroid hormone receptor, said method comprising:
- 4 (a) contacting the polypeptide of claim 23 with
- 5 a parathyroid hormone, (i) in the presence or (ii) in the
- 6 absence of a candidate compound; and
- 7 (b) comparing (i) the level of binding of said
- 8 polypeptide to said parathyroid hormone in the presence
- 9 of said candidate compound, with (ii) the level of
- 10 binding of said polypeptide to said parathyroid hormone
- 11 in the absence of said candidate compound; a lower level
- 12 of binding in the presence of said candidate compound
- 13 than in its absence indicating that said candidate
- 14 compound is capable of competing with said parathyroid
- 15 hormone for binding to said receptor.
 - 1 32. A method for identifying a compound capable
- 2 of competing with a parathyroid hormone-related protein
- 3 for binding to a parathyroid hormone receptor, said
- 4 method comprising:
- 5 (a) contacting the polypeptide of claim 23 with
- 6 a parathyroid hormone-related protein, (i) in the
- 7 presence or (ii) in the absence of a candidate compound;
- 8 and
- 9 (b) comparing (i) the level of binding of said
- 10 polypeptide to said parathyroid hormone-related protein
- in the presence of said candidate compound, with (ii) the
- 12 level of binding of said polypeptide to said parathyroid
- 13 hormone-related protein in the absence of said candidate
- 14 compound; a lower level of binding in the presence of
- 15 said candidate compound than in its absence indicating
- 16 that said candidate compound is capable of competing with

- said parathyroid hormone-related protein for binding to said receptor.
- 1 33. A method for identifying a compound capable of competing with a parathyroid hormone for binding to a
- 3 parathyroid hormone receptor, said method comprising:
- 4 (a) combining a parathyroid hormone with the
- 5 cell of claim 11, (i) in the presence or (ii) in the
- 6 absence of a candidate compound; and
- 7 (b) comparing (i) the level of binding of said
- 8 receptor to said parathyroid hormone in the presence of
- 9 said candidate compound, with (ii) the level of binding
- 10 of said receptor to said parathyroid hormone in the
- 11 absence of said candidate compound; a lower level of
- 12 binding in the presence of said candidate compound than
- in its absence indicating that said candidate compound is
- 14 capable of competing with said parathyroid hormone for
- 15 binding to said receptor.
 - 1 34. A compound capable of inhibiting the binding
 - 2 of parathyroid hormone or parathyroid hormone-related
 - 3 protein to a parathyroid receptor on the surface of a
- 4 cell.
- 1 35. A therapeutic composition comprising the
- 2 compound of claim 34 and a pharmaceutically-acceptable
- 3 carrier.
- 1 36. A method for identifying a DNA sequence
- 2 homologous to a parathyroid hormone receptor-encoding DNA
- 3 sequence, said method comprising:
- 4 providing a genomic or cDNA library;
- 5 contacting said library with the single-
- 6 stranded DNA of claim 18, under conditions permitting

- 7 hybridization between said single-stranded DNA and a
- 8 homologous DNA sequence in said library; and
- 9 identifying a clone from said library which
- 10 hybridizes to said single-stranded DNA, said
- 11 hybridization being indicative of the presence in said
- 12 clone of a DNA sequence homologous to a parathyroid
- 13 hormone receptor-encoding DNA sequence.
- 1 37. A transgenic non-human vertebrate animal
- 2 bearing a transgene comprising a DNA sequence encoding
- 3 parathyroid hormone receptor or a fragment thereof.
- 38. A diagnostic method comprising:
- 2 (a) obtaining a first blood sample from an
- 3 animal; (b) administering the composition of claim
- 4 35 to said animal;
- 5 (c) obtaining a second blood sample from said
- 6 animal subsequent to said administration of said
- 7 composition; and
- 8 (d) comparing the calcium level in said first
- 9 blood sample with that in said second blood sample, a
- 10 lower calcium level in said second blood sample being
- 11 diagnostic for a parathyroid hormone-related condition.
- 12 39. The isolated DNA of claim 1, wherein said
- 13 DNA sequence encodes a parathyroid hormone receptor.

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- 2 40. The parathyroid hormone receptor of claim 20
- 3 for use in therapy or diagnosis.
- 4 41. The polypeptide of claim 23 for use in
- 5 therapy or diagnosis.
- 6 42. The antibody of claim 27 for use in therapy
- 7 or diagnosis.

Mary.

- 8 43. The therapeutic composition of claim 26 for
- 9 use in therapy for the inhibition of activation by
- 10 parathyroid hormone or parathyroid hormone-related
- 11 protein of a parathyroid hormone receptor of a mammal or
- 12 for the reduction of the level of calcium in the blood of
- 13 a mammal.

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- 14 44. The therapeutic composition of claim 28 for
- 15 use in therapy for the inhibition of activation by
- 16 parathyroid hormone or parathyroid hormone-related
- 17 protein of a parathyroid hormone receptor of a mammal or
- 18 for the reduction of the level of calcium in the blood of
- 19 a mammal.
- 20 45. The parathyroid hormone receptor of claim 20
- 21 for use in the manufacture of a medicament for use in
- 22 therapy for the inhibition of activation by parathyroid
- 23 hormone or parathyroid hormone-related protein of a
- 24 parathyroid hormone receptor of a mammal or for the
- 25 reduction of the level of calcium in the blood of a
- 26 mammal.
- 27 46. The polypeptide of claim 23 for use in the
- 28 manufacture of a medicament for use in therapy for the
- 29 inhibition of activation by parathyroid hormone or
- 30 parathyroid hormone-related protein of a parathyroid
- 31 hormone receptor of a mammal or for the reduction of the
- 32 level of calcium in the blood of a mammal.
- 33 47. The antibody of claim 27 for use in the
- 34 manufacture of a medicament for use in therapy for the
- 35 inhibition of activation by parathyroid hormone or
- 36 parathyroid hormone-related protein of a parathyroid
- 37 hormone receptor of a mammal or for the reduction of the
- 38 level of calcium in the blood of a mammal.

- 48. A method for identifying a hypercalcemic condition in a patient which is mediated by parathyroid hormone or parathyroid hormone-related protein, the
- 42 method comprising
- (a) determining the calcium level of a first blood sample from the patient,
- (b) determining the calcium level of a second blood sample from the patient taken at a time subsequent after administration of the therapeutic composition of claim 26, and
- (c) comparing the calcium levels of the two
 blood samples, a lower calcium level in the second blood
 sample being indicative of a condition related to
 parathyroid hormone or parathyroid hormone-related
 protein in the patient.
- 49. A method for identifying a hypercalcemic condition in a patient which is mediated by parathyroid hormone or parathyroid hormone-related protein, the method comprising
- 58 (a) determining the calcium level of a first 59 blood sample from the patient,
- (b) determining the calcium level of a second blood sample from the patient taken at a subsequent time after administration of the therapeutic composition of claim 28, and
- (c) comparing the calcium levels of the two blood samples, a lower calcium level in the second blood sample being indicative of a condition related to parathyroid hormone of parathyroid hormone-related protein in the patient.

FIG. 1

•	TGGC	TGGGCACAGC CACCCTGTTG GTAGTCCAGG GGCCAGCCCA CTGAGCTGGC ATATCAGCTG													δŨ		
r	GTGG	GTGGCCCCGT TGGACTCGGC CCTAGGGAAC GGCGGCG ATG GGA GCG CCC CGG ATC Met Gly Ala Pro Arg Ile 1 5															115
	TCG Ser	CAC His	AGC Ser	CTT Leu 10	GCC Ala	TTG Leu	CTC Leu	CTC Leu	TGC Cys 15	TGC Cys	TCC Ser	GTG Val	CTC Leu	AGC Ser 20	TCC Ser	GTC Val	157
	TAC Tyr	GCA Ala	CTG Leu 25	GTG Val	GAT Asp	GCC Ala	GAT Asp	GAT Asp 30	GTC Val	λΤΆ Ile	ACG Thr	AAG Lys	GAG Glu 35	GAG Glu	CAG Gln	ATC Ile	2:-
	ATT Ile	CTT Leu 40	CTG Leu	CGC Arg	AAT Asn	GCC Ala	CAG Gln 45	GCC Ala	CAG Gln	TG T Cys	GAG Glu	CAG Gln 50	CGC Arg	CTG Leu	AAA Lys	GAG Glu	259
	GTC Val 55	CTC Leu	AGG Arg	GTC Val	CCT Pro	GAA Glu 60	CTT Leu	GCT Ala	GAA Glu	TCT Ser	GCC Ala 65	AAA Lys	GAC Asp	TGG Trp	ATG Met	TCA Ser 70	307
	AGG Arg	TCT Ser	GCA Ala	AAG Lys	ACA Thr 75	AAG Lys	AAG Lys	GAG Glu	AAA Lys	CCT Pro 80	GCA Ala	GAA Glu	AAG Lys	CTT Leu	TAT Tyr 85	CCC Pro	355
	CAG Gln	GCA Ala	GAG Glu	GAG Glu 90	TCC Ser	AGG Arg	GAA Glu	GTT Val	TCT Ser 35	GAC Asp	AGG Arg	AGC Ser	CGG Arg	CTG Leu 100	Gln	GAT Asp	403
	3GC 3ly	TTC Phe	TGC Cys 105	CTA Leu	CCT Fro	GA G Glu	125 136	ASP	AAC Asn	ATT	GTG Val	TGC Cys	TIP TIP	CCT Pro	GCT Ala	G GA Gly	4: _
	GTG Val	CCC Pro 120	GGC Gly	AAG Lys	GTG Val	GTG Val	GCC Ala 125	GTG Vai	SCC	TGC Cys	ccc	GAC Asp 130	TAC Tyr	TTC Phe	TAC Tyr	GAC Asp	499
	TTC Phe 135	AAC Asn	CAC His	A AA Lys	GGC	CGA Arg 140	Ala	TAT	CGG Arg	CGC	TGT Cys 145	GAC Asp	AGC Ser	AAT Asn	G GC Gly	AGC Ser 150	547
•	TGG Trp	GAG Glu	CTG Leu	GTG Val	CCT Pro 155	Gly	AAC Asn	AAC Asn	Arg CG G	ACA Thr 160	TGG	GCG Ala	AAT Asn	TAC Tyr	AGC Ser 165	GAA Glu	595
	TGT Cys	GTC Val	AAG Lys	TTT Phe 170	Leu	ACC	AAC Asn	GAG Glu	ACC Thr 175	Arg	GAA Glu) Arg	GAA Glu	GTC Val 180	TTT	GAT Asp	643

FIG. :

J		185			1-	1111	190	GTÅ	TYE	. Sei	- II	e Se	r Le 5	eu G	C TCC y Ser	
	200				. 500	205	, Dea	GIY	- 2 -	rne	21	g Ar	g Le	u Hí	T TGC s Cys	739
215			-1-	1,10	220	Mec	nis	Leu	Sue	225	Sei	r Pho	e Me	t Le	C CGG u Arg 230	787
			416	235	116	пåз	vañ	via	240	Leu	туг	Sex	G1	y Va 24		835
			250	014	ur A	115	1414	235	GIU	GIU	Leu	Arg	Ala 260	Pho	ACA Thr	883
		265		u T G	vañ	LYS	270	arā	LUG	Val	Gly	Cys 275	Arç	[Va]		931
	2,80			Ted	- 71	285	red	-nr	Thr	ASN	777 290	Tyr	Trp	Įle		979 ·
GTG (Val (295		- 1		-1-	300	UIS	ser	_au	TIE	305	Met	Ala	Phe	Phe	Ser 310	1027
GAG A	-, -	-,3	-1-	315	ırp	GI?	3		TTA Leu 120	TTT Fhe	GGC Gly	TGG Trp	GGC Gly	CTC Leu 325	CCT Pro	1075
Ala :	al:	?he	7a1 330	Ala		775	`*a_									1123
ACT G	3	45			Leu .	SEL :	250	- <u>(</u>	sn .	Lys :	Lys	Trp 355	Ile	Ile	Gln	1171
	60					365	aı.	al A	sn i	rne :	11e 370	Leu	Phe	Ile	Asn	1219
ATA A' Ile I: 375	TC A le A	.GA (GTC (GCT A Ala 1 380	ACT 3 Thr I	AAA C	TC C A us.	rd (GAG A Glu 1 885	ACC .	AAT (Asn)	GCA Ala	G GG Gly	AGA Arg 390	1267

TG:	GAC Asp	ACG Thr	AGG Arg	CAA Gln 395	CAG Gln	TAT Tyr	AGA Arg	AAG Lys	CTG Leu 400	CTG Leu	lys	TCC Ser	ACG Thr	CTA Leu 405	GTC Val	131
CTC	ATG Met	CCG Pro	CTA Leu 410	TTT	GCG	GTG Val	CAC His	TAC Tyr 415	ATC Ile	GTC Val	TTC Phe	ATG Met	GCC Ala 420	ACG Thr	CCG Pro	1353
TAC Tyr	ACA Thr	GAA Glu 425	GTA Val	TCA Ser	GGG Gly	ATT Ile	CTT Leu 430	TGG Trp	CAA Gln	GTC Val	CAA Gln	ATG Met 435	CAC	TAT Tyr	GAA Glu	1411
ATG Met	CTC Leu 440	TTC Phe	AAT Asn	TCA Ser	TTC Phe	CAG Gln 445	GGA Gly	TTT Phe	TTC Phe	GTT Val	GCC Ala 450	ATT Ile	ATA Ile	TAC Tyr	TGT Cys	1453
TTC Phe 455	TGC Cys	AAT Asn	GGA Gly	GAG Glu	GTA Val 460	CAA Gln	GCA Ala	GAG Glu	ATC Ile	AAG Lys 465	AAG Lys	TCA Ser	TGG Trp	AGC Ser	CGA Arg 470	1507
TGG Trp	ACC Thr	CTG Leu	GCC Ala	TTG Leu 475	GAC Asp	TTC Phe	AAG Lys	cgg	AAG Lys 480	GCC Ala	CGG Arg	AGT Ser	GGC Gly	AGC Ser 485	AGT Ser	1555
ACC	TAC Tyr	AGC Ser	TAT Tyr 490	GGC	CCC Pro	ATG Met	GTG Val	TCA Ser 495	CAT His	ACA Thr	AGT Ser	Val	ACC Thr 500	AAT Asn	GTG Val	1603
GGA Gly	CCT Pro	CGA Arg 505	GGG Gly	GGC Gly	TGG Trp	CCT Pro	TGT Cys 510	CCC Pro	TCA Ser	GCC Ala	Leu	GAC Asp 515	TAGC	TCCT	GG	1652
GGC	rggag	CC A	GTGC	CAAT	'G GC	CATO	ACCA	GTI	cci	'GGC	TATG	TGAA	GC A	TGGT	TCCAT	1712
TTC	Gaga	AC I	CATI	GCCT	T CA	TCTG	GCCC	: AGA	cct	'GGC	ACCA	AAGA'	TG A	CGGG	TATCT	1771
CAA	GGCT	cr c	GACT	TTAT	G YC	CCAA	TGGT	TGG	GGAA	.CAG	cccc	ctcc	ac t	ccrs	GAGGA	1832
GGAC	AGAG	AG A	CAGT	CATG	T GA	CCCA	TATO	:								1862

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FIG. 2

TGG	GCAC	AGC	CACC	CTGT	TG G	TAGT	CCAG	G GG	CCAG	CCCA	CTC	AGCI	rggc	ATA	CAGCTG	60
GTG	GCCC	CGT	TGGA	CTCG	GC C	CTAG	GGAA	c GG	CGGC	G AT Me	G GG t Gl 1	A GC Y Al	G CC a Pr	C CC	G ATC g Ile 5	115
TCG Ser	CAC His	AGC Ser	CTT Leu 10	GCC Ala	TTG Leu	CTC Leu	CTC Leu	TGC Cys 15	TGC Cys	TCC Ser	GTG Val	CTC	AGC Ser 20	TCC Ser	GTC Val	163
TAC Tyr	GCA Ala	CTG Leu 25	GTG Val	G AT Asp	GCC Ala	GAT Asp	GAT ASD 30	GTC Val	ATA Ile	ACG Thr	AAG Lys	GAG Glu 35	G AG Glu	CAG Gln	ATC	211
ATT	CTT Leu 40	CTG Leu	CGC Arg	AAT Asn	GCC Ala	CAG Gln 45	GCC Ala	CAG Gln	TGT Cys	GAG Glu	CAG Gln 50	CGC Arg	CTG Leu	AAA Lys	GAG Glu	259
GTC Val 55	CTC Leu	AGG Arg	GTC Val	CCT Pro	GAA Glu 60	CTT Leu	GCT Ala	GAA Glu	TCT Ser	GCC Ala 65	AAA Lys	GAC Asp	TGG Trp	ATG Met	TCA Ser 70	307
AGG Arg	TCT Ser	GCA Ala	AAG Lys	ACA Thr 75	AAG Lys	AAG Lys	GAG Glu	λλλ Lys	CCT Pro 80	GCA Ala	GAA Glu	AAG Lys	CTT Leu	TAT Tyr 85	CCC Pro	355
CAG Gln	GCA Ala	GAG Glu	GAG Glu 90	TCC Ser	AGG Arg	GAA Glu	GTT Vai	TCT Ser	GAC Asp	AGG Arg	AGC Ser	CGG Arg	CTG Leu 100	Gln	GAT Asp	403
GGC 317	TTC Fhe	TGC Cys 105	CTA Lau	CCT Pro	GAG Glu	TGG	GAC ET	AAC .ET.	ATT	GTG Val	TGC Cys	TGG Trp 115	CCT Pro	GCT Ala	GGA Gly	451
GTG Val	CCC Pro 120	GGC Gly	AAG Lys	GTG Val	GTG Val	GCC Ala 125	373	000	TGC Cys	CCC Pro	GAC Asp 130	TAC Tyr	TTC Phe	TAC Tyr	GAC Asp	499
TTC Phe 135	AAC Asn	CAC His	AAA Lys	GGC Gly	CGA Arg 140	GCC Ala	TAT Tyr	ogg Arg	CGC Arg	TGT Cys 145	GAC Asp	AGC Ser	AAT Asn	GGC Gly	AGC Ser 150	547
TGG Trp	GAG Glu	CTG Leu	GTG Val	CCT Pro 155	GGG Gly	AAC Asn	AAC Asn	e gg	ACA Thr 160	TGG Trp	GCG Ala	AAT Asn	TAC Tyr	AGC Ser 165	GAA Glu	595
TGT Cys	GTC Val	AAG Lys	TTT Phe 170	CTG Leu	ACC Thr	AAC Asn	GAG Glu	ACC The 175	CGG Arg	G AA Glu	CGG Arg	GAA Glu	GTC Val 180	TTT Phe	GAT Asp	643

FIG. 2

CGC CTC GGA ATG ATC TAC ACT GTG GGC TAC TCC ATC TCT CTG GGC TCC 591 Arg Leu Gly Met Ile Tyr Thr Val Gly Tyr Ser Ile Ser Leu Gly Ser 185 CTC ACT GTG GCT GTG CTG ATT CTG GGT TAC TTT AGG AGG TTA CAT TGC Leu Thr Val Ala Val Leu Ile Leu Gly Tyr Phe Arg Arg Leu His Cys 200 205 ACC CGA AAC TAC ATT CAC ATG CAT CTC TTC GTG TCC TTT ATG CTC CGG Thr Arg Asn Tyr Ile His Met His Leu Phe Val Ser Phe Met Leu Arg 215 GCT GTA AGC ATC TTC ATC AAG GAT GCT GTG CTC TAC TCG GGG GTT TCC 335 Ala Val Ser Ile Phe Ile Lys Asp Ala Val Leu Tyr Ser Gly Val Ser 235 240 ACA GAT GAA ATC GAG CGC ATC ACC GAG GAG GAG CTG AGG GCC TTC ACA Thr Asp Glu Ile Glu Arg Ile Thr Glu Glu Glu Leu Arg Ala Phe Thr 383 250 255 GAG CCT CCC CCT GCT GAC AAG GCG GGT TTT GTG GGC TGC AGA GTG GCG 931 Glu Pro Pro Pro Ala Asp Lys Ala Gly Phe Val Gly Cys Arg Val Ala 265 270 GTA ACC GTC TTC CTT TAC TTC CTG ACC ACC AAC TAC TAC TGG ATC CTG 979 Val Thr Val Phe Leu Tyr Phe Leu Thr Thr Asn Tyr Tyr Trp Ile Leu 280 285 GTG GAA GGC CTC TAC CTT CAC AGC CTC ATC TTC ATG GCT TTT TTC TCT 1027 Val Glu Gly Leu Tyr Leu His Ser Leu Ile Phe Met Ala Phe Phe Ser 295 300 305 GAG AAA AAG TAT CTC TGG GGT TTC ACA TTA TTT GGC TGG GGC CTC CCT 1073 Glu Lys Lys Tyr Leu Trp Gly Phe Thr Leu Phe Gly Trp Gly Leu Pro GCC GTG TTT GTC GCT GTG TGG GTG ACC GTG AGG GCT ACA CTG GCC AAC 1120 Ala Val Phe Val Ala Val Trp Val Thr Val Arg Ala Thr Leu Ala Asn 330 335 ACT GAG TGC TGG GAC CTG AGT TCG GGG AAT AAG AAA TGG ATC ATA CAG 1171 Thr Glu Cys Trp Asp Leu Ser Ser Gly Asn Lys Lys Trp Ile Ile Gln 345 GTG CCC ATC CTG GCA GCT ATT GTG GTG AAC TTT ATT CTT TTT ATC AAT 1219 Val Pro Ile Leu Ala Ala Ile Val Val Asn Phe Ile Leu Phe Ile Asn 360 ATA ATC AGA GTC CTG GCT ACT AAA CTC CGG GAG ACC AAT GCA GGG AGA 1267 Ile Ile Arg Val Leu Ala Thr Lys Leu Arg Glu Thr Asn Ala Gly Arg 375 380 385

FIG. 2

	AGG CAA CAG TAT Arg Gln Gln Tyr 395				1315
	CTA TTT GGG GTG Leu Phe Gly Val 410				1363
	GTA TCA GGG ATT Val Ser Gly Ile				1411
	AAT TCA TTC CAG Asn Ser Phe Gln 445				1459
	GGA GAG GTA CAA Gly Glu Val Gln 460				1507
	GCC TTG GAC TTC Ala Leu Asp Phe 475				1555
	TAT GGC CCC ATG Tyr Gly Pro Met 490				1603
	GGG GGG CTG GCC Gly Gly Leu Ala				1651
	GCC AGT GCC AAT Ala Ser Ala Asn 525				1699
Lys His Gly	Ser Ile Ser Glu 540	Asm Ser Leu	Pro Ser Ser		1747
	AAA GAT GAC GGG Lys Asp Asp Gly 555		Gly Ser Gly		1795
CCA ATG GTT Pro Met Val	GGG GAA CAG CCC Gly Glu Gln Pro 570	CCT CCA CTC Pro Fro Leu 575	CTG GAG GAG Leu Glu Glu	GAG AGA GAG Glu Arg Glu 580	1843
ACA GTC ATG Thr Val Met 585	TGACCCATAT C				1863

FIG. 3

GG	CGGGG	GCC _.	GCGG	CGGC	GA G	CTCG	GAGG	C CG	GCGG	CGGC	TGC	ccc	AGG	GAC	ceeccc	60
TAC	GCGG	TGG	CG A	TG G et G 1	GG G ly A	CC G la A	CC C la A	GG A rg I 5	TC G le A	CA C la P	cc A	GC C er L	TG o eu A 10	SCG (TC .eu	108
CTA	CTC Leu	TGC Cys 15	TGC Cys	CCA Pro	GTG Val	crc Leu	AGC Ser 20	TCC Ser	GCA Ala	TAT	GCG Ala	CTG Leu 25	GTG Val	GAT Asp	GCG Ala	133
GAC Asp	GAT Asp 30	Val	TTT Phe	ACC Thr	AAA Lys	GAG Glu 35	GAA Glu	CAG Gln	ATT Ile	TTC Phe	CTG Leu 40	CTG Leu	CAC His	CGT Arg	GCC Ala	204
CAG Gln 45	GCG Ala	CAA Gln	TGT Cys	GAC Asp	AAG Lys 50	CTG Leu	CTC Leu	λAG Lys	GAA Glu	GTT Val 55	CTG Leu	CAC His	ACA Thr	GCA Ala	GCC Ala 60	27-
AAC Asn	ATA Ile	ATG Met	GAG Glu	TCA Ser 65	GAC Asp	AAG Lys	GGC Gly	TGG Trp	λCA Thr 70	CCA Pro	GCA Ala	TCT Ser	ACG Thr	TCA Ser 75	GG G Gly	300
AAG Lys	CCC Pro	AGG Arg	AAA Lys 80	GAG Glu	AAG Lys	GCA Ala	TCG Ser	GGA Gly 85	AAG Lys	TTC Phe	TAC Tyr	CCT Pro	GAG Glu 90	TCT Ser	A AA Lys	348
GAG Glu	AAC Asn	AAG Lys 95	GAC Asp	GTG Val	CCC Pro	ACC Thr	GGC Gly 100	Ser	AGG Arg	CGC Arg	AGA Arg	GGG Gly 105	Arg	CCC Pro	TGT Cys	396
CTG Leu	CCC Pro 110	GAG Glu	TGG Trp	GAC Asp	AAC Asn	ATC Ile	GTT Val	TGC Cys	TGG	CCA Pro	TTA Leu 120	GGG Gly	GCA Ala	CCA Pro	GGT Gly	444
GAA 31:1 125	GTG Val	GTG Tal	GCA Ala	GTA Tal	CCT Pro 130	TGT	ccc Pro	GAT Asp	TAC Tyr	ATT Ile ISS	TAT	GAC Asp	TTC Phe	AAT Asn	CAC His 140	490
AAA Lys	GGC Gly	CAT His	GCC Ala	TAC Tyr 145	AGA Arg	CGC Arg	TGT Cys	GAC Asp	CGC Arg 150	AAT Asn	GGC Gly	λGC Ser	TGG Trp	GAG Glu 155	GTG Val	54C
	CCA Pro															588
TTC Phe	ATG Met	ACC Thr 175	AAT Asn	GAG Glu	ACG Thr	cgg Arg	GAA Glu 180	cgg Arg	GAG Glu	GTA Val	TTT Phe	GAC Asp 185	CGC Arg	CTA Leu	GGC Gly	636
	ATC Ile 190															684

FIG. 3

20	5		4 T.	- reu	210	Tyr	Pne	AFG	Arg	215	1 HT	s Cys	Thi	Arq	C AAC J Asn 220	732	
*11		- 1115	nec	225	met	aue	Leu	ser	230	Met	Lei	ı Arq	Ala	235		780	
110	FILE	, val	240	Asp	Ala	Val	Leu	245	ser	GIY	Phe	? Thr	Leu 250	Asp		828	
VIG	. 014	255	Leu	inr	GIU	GIU	250	Lau	nls	Ile	Ile	GCG Ala 265	Gln	Val	Pro	876	
11,0	270		AIG	Ala	AIA	275	vaı	CTÅ	ryr	Ala	Gly 280		Arg	Val	Ala	924	
285		rne	FIIĢ	290	TYT	Pne	reu	YIG	7nr 295	ASN	Tyr	TAC	Trp	Ile 300	Leu	972	
, 41	914	Gly	305	-yr	Leu	nıs	ser	310	lle	Phe	Met	GCC Ala	Phe 315	Phe	Ser	1020	
J1 4	Lys	320	TYP	Leu	Trp	GIĀ	325	Thr	ile	Phe	Gly	TGG Trp 330	Gly	Leu	Pro	1068	
	335	2116	·41	A12	AT	340	∨a1	3 <u>2 '</u> 4	∵al	Arg	Ala 345	ACC Thr	Leu	Ala	Asa	1116	
ACT Thr 350	GGG Gly	TGC Cys	Tog	G AT Asp	crg Leu 355	AGC Ser	TCC Ser	133 117	:15	AAG Lys 360	Lys	TGG Trp	ATC Ile	Ile	CAG Gln 365	1164	
GTG Val	CCC Pro	ATC Ile	CTG Leu	GCA Ala 370	TCT Ser	GTT Val	GTG Val	_au	AAC Asn 375	TTC . Phe	ATC Ile	CTT :	Phe	ATC Ile 380	AAC Asn	1212	
ATC	ATC Ile	wr.a	GTG Val 385	CTT Leu	GCC Ala	ACT . Thr	Lys	CTT Leu 190	ccc Arg	GAG /	ACC Thr	Asn .	GCG Ala 395	GGC Gly	CGG Arg	1260	
TG T Cys	vañ	ACC Thr 400	AGG Arg	CAG Gln	CAG '	TYT .	CGG . Arg 405	rys .	CTG (Leu)	CTC /	Arg	TCC : Ser :	ACG '	TTG (Leu '	GTG Val	1308	

CTC GTG CC Leu Val Pr 415	G CTC TTT GG o Leu Phe Gl	GTC CAC T Val His T 420	AC ACC GTC	TTC ATG GCC TTG Phe Met Ala Leu 425	CCG 1356 Pro
TAC ACC GA Tyr Thr G1 430	G GTC TCA GGC u Val Ser Gly 435	Thr Leu T	GG CAG ATC rp Gln Ile 440	CAG ATG CAT TAT Gln Met His Tyr	GAG 1404 Glu 445
ATG CTC TT Met Leu Ph	C AAC TCC TTC e Asn Ser Phe 450	CAG GGA T	TT TTT GTT he Phe Val 455	GCC ATC ATA TAC Ala Ile Ile Tyr (IGT 1452 Cys
TTC TGC AA Phe Cys As:	r GGT GAG GTG n Gly Glu Val 465	GIN Ala G.	AG ATT AGG : lu Ile Arg : 70	AAG TCA TGG AGC (Lys Ser Trp Ser) 175	CGC 1500 Arg
TGG ACA CTO Trp Thr Lev 480	ı ala Len Asb	TTC AAG CO Phe Lys Ai 485	GC AAA GCA (rg Lys Ala A	CGA AGT GGG AGT A Arg Ser Gly Ser S 490	AGC 1548 Ser
AGC TAC AGG Ser Tyr Sei 495	TAT GGC CCA Tyr Gly Pro	ATG GTG TO Met Val Se 500	er His Thr S	AGT GTG ACC AAT G Ser Val Thr Asn V 505	TG 1596
GGC CCC CGT Gly Pro Arc 510	GCA GGA CTC Ala Gly Leu 515	AGC CTC CC Ser Leu Pr	CC CTC AGC C TO Leu Ser F 520	CCC CGC CTG CCT C Pro Arg Leu Pro P 5	CT 1644 ro 25
GCC ACT ACC Ala Thr Thr	AAT GGC CAC Asn Gly His 530	TCC CAG CT Ser Gln Le	CG CCT GGC C EU Pro Gly H 535	CAT GCC AAG CCA G His Ala Lys Pro G 540	GG 1692 ly
AIA PRO AIA	Thr Glu Thr	Glu Thr La	u Pro Val T	ACT ATG GCG GTT C Thr Met Ala Val P 555	ro
AAG GAC GAT Lys Asp Asp 560	Gly Fhe Lau	AAC SGC TO Asn Gly Se 565	C TGC TCA G	GC CTG GAT GAG G ly Leu Asp Glu G 570	AG 1783 lu
GCC TCC GGG Ala Ser Gly 575	TCT GCG CGG Ser Ala Arg	CCG CCT CC Pro Pro Pr 580	o Leu Leu G	AG GAA GGA TGG G In Glu Gly Trp G 85	AA 1836 lu
ACA GTC ATG Thr Val Met 590	TGACTGGGCA (CTAGGGGGCT .	AGACTGCTGG	CCTGGGCACA 1885	
				ATTCGGGAT CTGGACC	
				AAGGAAGAG GTTTTGC	AGG 2005
AATTAAATAT	GITTCCTCAG T	A DTADTADD:	GGACACAAG G	AAGGC	2051

Fig. 4

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1 MGAARIAPSLALLLCCPVLSSAYALVDADDVFTREEQIFLLHRAQAQCDX 50
     1 MGAPRISHSLALLLCCSVLSSVYALVDADDVITKEEQIILLRNAQAQCEQ 50
  51 LLKEVLHTAANIHESDKGWTPASTSGKPRKEKASGKFYPESKENKDVPTG 100
      51 RIKEVIR. VPELAESAKOW. . HSRSAKTKKEKPAEKLYPQAEESREVSDR 97
 101 SRRRGRPCL?EWDNIVCWPLGAPGEVVAVPCPDYIYDFNHRGHAYRRCDR 150
     11 .: 11111111111 1.11.111111111:111111:111111.
  98 SRLODGFCLPENDNIVCWPAGVPGRVVAVPCPDYFYDFNHKGRAYRRCDS 147
 151 NGSWEVVPGHNRTWANYSECLAFHTNETREREVFDRLGHIYTVGYSHSLA 200
    148 NGSWELVPGNNRTWANYSECVRFLTNETREREVFDRLGHIYTVGYSISLG 197
 201 SLTYAVLILAYFRRLHCTRNYIEMHMFLSFHLRAASIFVKDAVLYSGFTL 250
    198 SUTVAVLILGYFRRLHCTRNYIHHHLFVSFHLRAVSIFIKDAVLYSGVST 247
 251 DEAERLTEEELHIIAQVPPPPAAAAVGYAGCRVAVTFFLYFLATNYYWIL 300
    248 DEIERITEEELRAFTE...PPPADKAGFVGCZVAVTVFLYFLITNYYWIL 294
 301 VEGLYLHSLIFHAFFSEKKYLWGFTIFGWGLFAVFVAVWVGVRATLANTG 350
    VEGLYLHSLIFMAFFSEKKYLWGFTLFGHGLFAVFVAVWVTVRATLANTE 344
351 CWDLSSGHKKWIIOVPILASVVLNFILFINIIRVLATKLRETNAGRCDTR 400
    345 CWDLSSGNKKWIIQVPILAAIVVNFILFINIIRVLATKLRETNAGRCDTR 394
401 OQYRKLLRSTLVLVPLFGVHYTVFMALPYTEVSGTLWQIQHHYEMLFNSF 450
   filitienineninen in inini-litenininini
395 QQYRKLLKSTLYLMPLFGVHYIVFMATPYTEVSGILWQVQHHYEHLFHSF 444
451 CGFFVAIIYCFCHGEVQAEIRKSWSRWTLALDFKRKARSGSSSYSYGPMV 500
   445 QGFFVAIIYCFCHGEVQAEIKKSWSRWTLALDFKRKARSGSSTYSYGPHV 494
501 SHTSVTNVGFPAGLSLPLSPRLPP...ATTNGHSQLPGHARPGAPATETE 547
   .... '111.....
495 SHTSYTTYGFRGGLALSLSPRLAFGAGASANGERQLFGYVKEGSISENSL 544
548 TLPVTMAVPKDDGFLNGSCSGLDEEASGSARPPPLLQEGWETVM. 591
545 PSSGPEPGTXDDGYLNG..SGLYEPMYG.EJPPPLLZEERETVM* 586
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Gap Weight: 3.000 Average Match: 0.540 Length Weight: 0.100 Average Mismatch: -0.396

Quality: 712.2 Length: 595
Ratio: 1.215 Gaps: 6
Percent Similarity: 87.113 Percent Edentity: 77.835
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Fig. 5

R15	MGAARIAPSL	ALLLCCPVLS	SAYALVDADD	VFTREEQIFE	LERAQAQCDE	50
Oko	MGAPRISHSL	ALLLCCSVLS	SVYALVDADD	VITREEQIIL	LRNAQAQCEQ	50
Okh				VITKEEQIIL	LRNAQAQCEQ	50
	-	A				
R15	LLKEVLHTAA	NIMESDKGWT	PASTSGKPRK	EXASGRETE	SKENKOVPTG	100
Oko	RLKEVLR. VP	ELAESAKDW.	. MSRSAKTKK	ERPAEKLYPO	AEESREVSOR	97
Okh					AZESREVSOR	
						•
	•	•		•	•	
R15 Oko	SRRRGRPCLP	EMDNIVCWPL	GAPGEVVAVP	CPDYIYDFNE	KGHAYRRCDR	150
Okh	SRIODGICLE	EMUNIVEWPA	CUPCEUVAUP	CPDYFYDFNR	KGRAYRRCDS KGRAYRRCDS	147
U	242004621		B		NOW INNEDS	147
	N	**	N			
R15	NGSWEVVPGH	NRTWANYSEC	LKFHTNETRE	REVFDRLGMI	YTVGYSHSLA	200
Oko	NGSWELVPGN	NRTWANYSEC	VKFLTNETRE	REVPDRLGHI	YTVGYSISLG	
Okh	NGSWELVPGN	NRTWANYSEC	VKFLTNETRE	REVFORLGHI		197

R15	SLTVAVLILA	YFRRLHCTRN	YIHMRMFLSF	MLRAASIFVK	DAVLYSGFTL	250
Oko	SLTVAVLILG	YFRRLHCTRN	YIHMHLFVSF	HLRAVSIFIK	DAVLYSGVST	247
Okh	SLTVAVLILG	YFRRLHCTRN	YIHHHLFVSF	MLRAVSIFIK	DAVLYSGVST	247
	C			D		
R15	DERED! #555	THTTSOUDDD	BALLAUGYAG		PLATNYYWIL	200
Oko	DEIERITEEE	LRAFTEP	PPADKAGEVG	CRVAVIVELY	FLTTNYYWIL	294
Okh					FLTTNYYWIL	
				E		
R15 Oko					GVRATLANTG TVRATLANTE	
Okh					TVRATLANTE	
				G		,,,
	•					
R15	CWDLSSGHKK	WIIQVPILAS	VVLNFILFIN	IIRVLATKLE	ETNAGRODTR	400
Oko					ETNAGRODTR	
Okh	CADT22CVKK		H		ETRAGREDTR	194
R15					REYEMLINSF	
Oko					xeyemlfnsf	
Okh	QQYRKLLKST			EAZCITMOAO	ABYEMLFNSF	444
		I				
R15	OGFFVALLYC	FCNGEVOAEI	RKSWSRWTLA	LDFKRKARSG	SSSYSYGPMV	500
Oko					SSTYSYGPHV	
Okh		-	KKSWSRWTLA	LDFKRKARSG	SSTYSYGPHV	494
	J					
R15	CHECHENICS	PACTOTOR	DI.DD 177	VGHSOT PGP1	KPGAPATETE	547
Oko					KHGSISENSL	
Okh	SHISVINVGP	RGG	%PCPSA	LD		515
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R15	TLPVTMAVPK	DDGFLNGSCS	GLDEEASGSA	REPELLQEGN	ETVA	591
Oko	rssgrergtx	DDGYLNGS	CLIEFANG.E	GEFFLLEEER	LIVA	585

FIG. 6

With 1 enzymes: SACI

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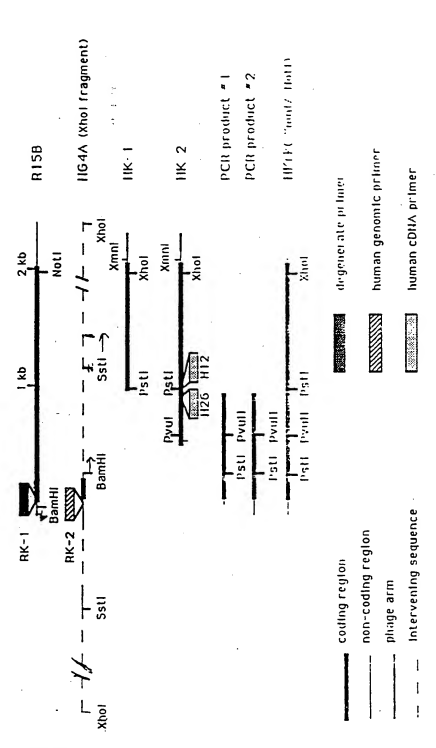
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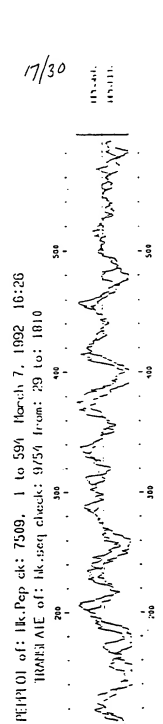
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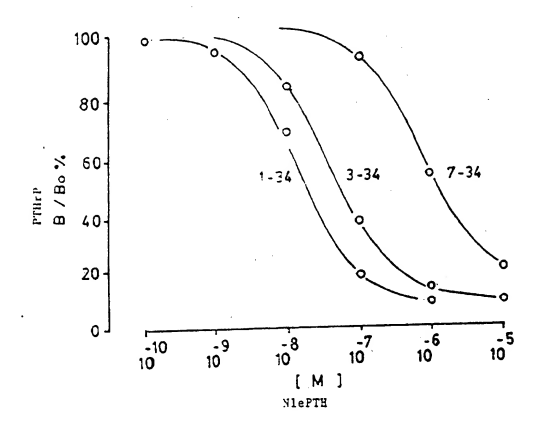


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Fig.3





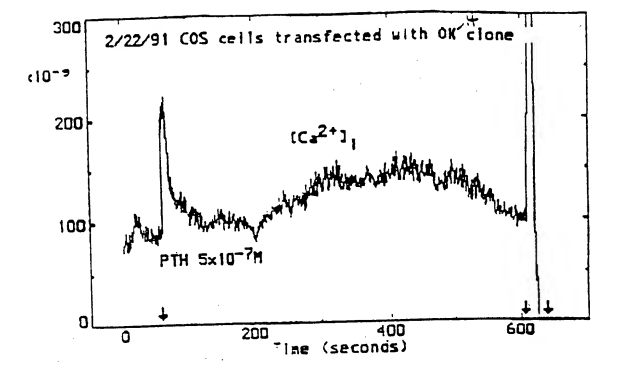
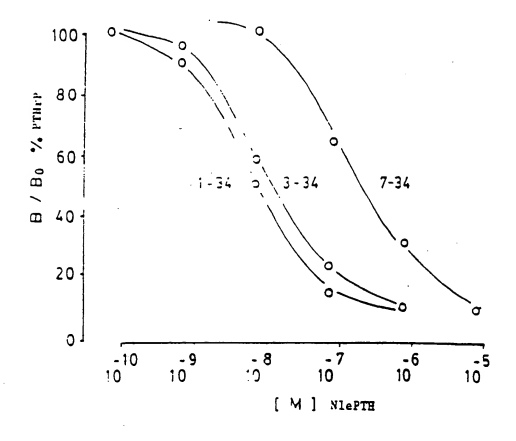
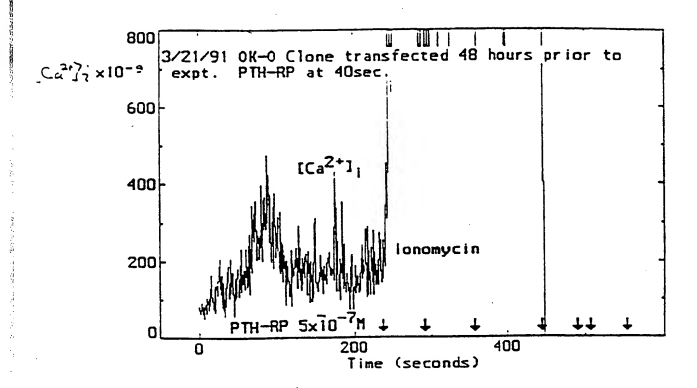


Fig. 11





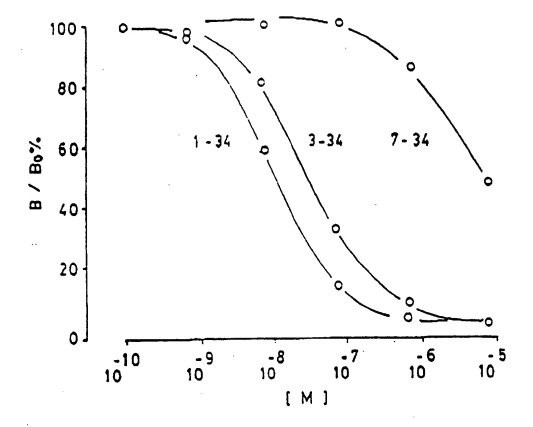
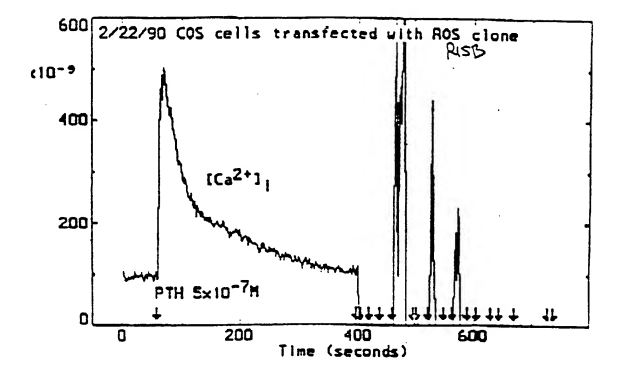
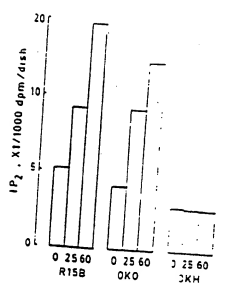
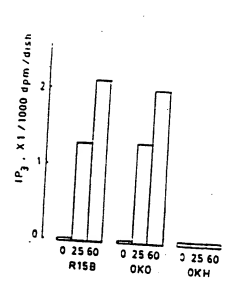
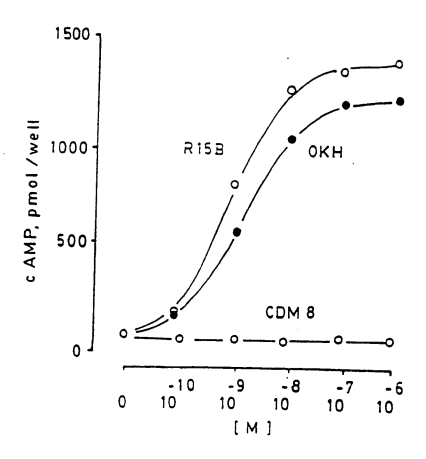


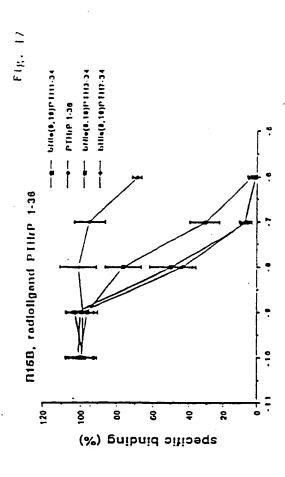
FIG. 13

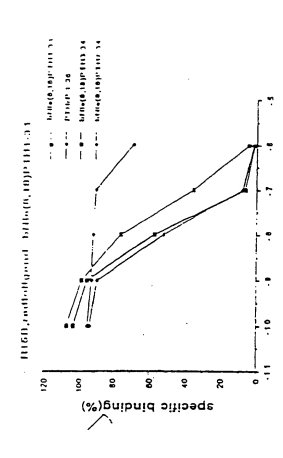


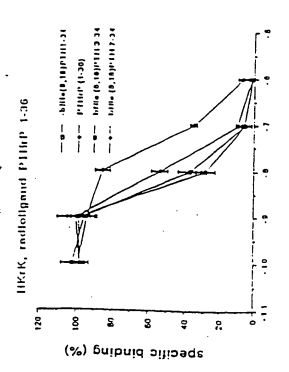


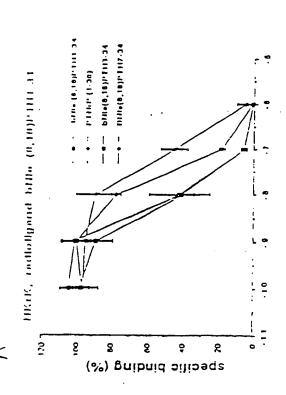






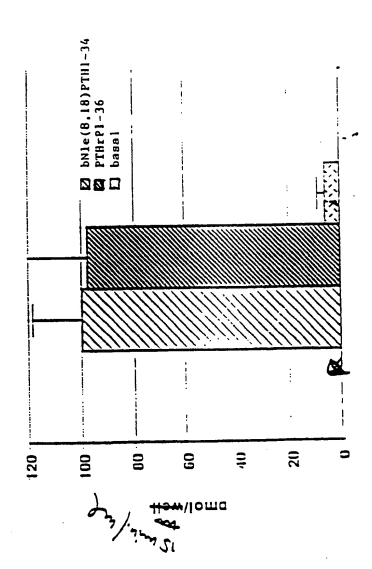






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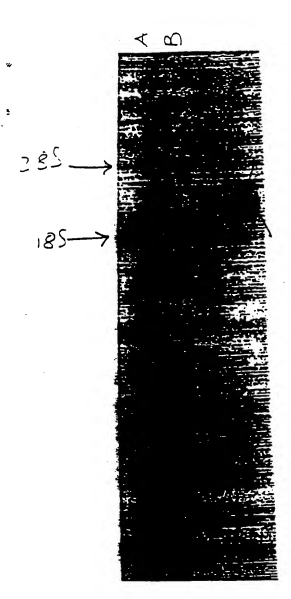
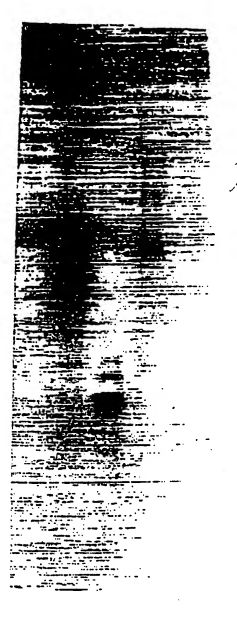
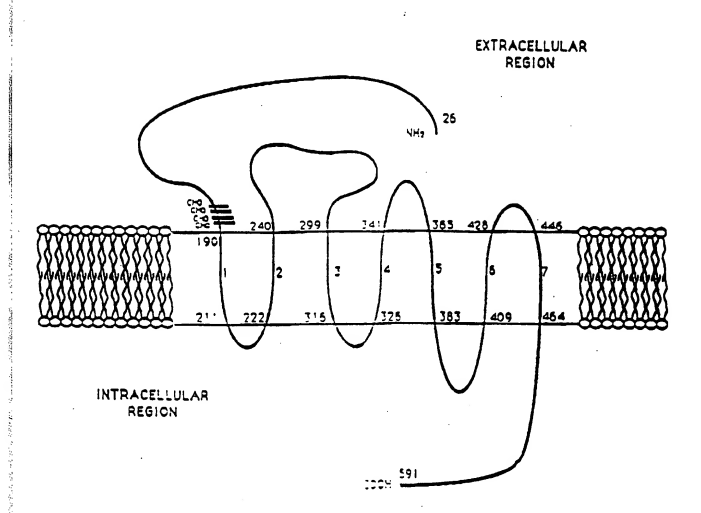


Fig. 19

Fig. 20



RAT BONE PTH/PTHrP RECEPTOR



AMING ACID SEQUENCE OF 7 PUTATIVE TRANS-MEMBRANE REGIONS

INTERNATIONAL SEARCH REPORT

International application No. PCT/US92/02821

A. CLA	SSIFICATION OF SUBJECT MATTER								
IPC(5) :Please See Extra Sheet.									
US CL :435/69.1, 240.2, 320.1; 536/27, 28, 29; 530/350, 387, 397, 399.									
	o International Patent Classification (IPC) or to both	national classification and IPC							
	DS SEARCHED								
1	ocumentation searched (classification system followed								
U.S. :	APS AND COMMERCIAL DATABASES (DIALOC	G) 435/69.1, 240.2, 320.1; 536/27, 28,	29						
Documental	ion searched other than minimum documentation to the	e extent that such documents are included	in the fields searched						
Flectronic d	ata hase consulted during the international search (na	me of data base and, where practicable	search terms used)						
Electronic data base consulted during the international search (name of data base and, where practicable, search terms used) DIALOG AND ONLINE SEQUENCE SEARCH									
C. DOCUMENTS CONSIDERED TO BE RELEVANT									
Category*	Citation of document, with indication, where ap	propriate, of the relevant passages	Relevant to claim No.						
<u>X</u> Y	TWENTY-SEVENTH ANNUAL MEETING OF THE BIOLOGY, VOLUME 105, NO. 4, PT. 2, ISSUED		<u>1-19, 39</u> 20-38, 40-49						
	AL., "MOLECULAR CLONING OF A PARAT	THYROID HORMONE RECEPTOR-	333, 13 11						
	RELATED MEMBRANE PROTEIN FROM N DOCUMENT.	MOUSE BONE CELLS, ENTIRE							
Y	THE JOURNAL OF BIOLOGICAL CHEMISTS		1-49						
	JANUARY 1990, ABOU-SAMRA ET AL., "O ACTIVE BIOTINYLATED PARATHYROID HOI								
	ENTIRE DOCUMENT.								
Y	BIOCHEMISTRY, VOLUME 29, NO. 30, ISSUE		1-49						
	"PREPARATION AND CHARACTERIZATION (N TYR-36)-PATHYROID HORMONE RELATED								
	AFFINITY, PARTIAL AGONIST HAVING HIG WITH ITS RECEPTOR ON ROS 17/2.8 CE								
	DOCUMENT.								
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Funt	ner documents are listed in the continuation of Box C	. See patent family annex.							
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Date of the	actual completion of the international search	Date of mailing of the international search report							
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INTERNATIONAL SEARCH REPORT

International application No. PCT/US92/02821

A. CLASSIFICATION OF SUBJECT MATTER: IPC (5):

C12P 21/06; C12N 5/00, 15/00; C07H 15/12, 17/00; C07K 3/00; A61K 35/14, 37/24, 37/36.

BOX II. OBSERVATIONS WHERE UNITY OF INVENTION WAS LACKING

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